Jan Delaval please

'Access DB#\_\_\_75681'

# SEARCH REQUEST FORM

Scientific and Technical Information Center

Sanua Bari - 11 9/12/2
Requester's Full Name: $\frac{SABIHA}{SABIHA} \frac{SA2I}{SABIHA} = \frac{7414I}{SABIHA} = \frac{9/13/62}{3.000}$
Art Unit: 1616 Phone Number 30 5-3910 Serial Number: 10 10 3 6 8 15  Mail Box and Bldg/Room Location: 2019 Results Format Preferred (circle) PAPER DISK E-MAIL
Mail Dox and Didg Room Docation Results Politiat Fletched (choic) 171 Exposor E-MAIL
If more than one search is submitted, please prioritize searches in order of need.
Please provide a detailed statement of the search topic, and describe as specifically as possible the subject matter to be searched.  Include the elected species or structures, keywords, synonyms, acronyms, and registry numbers, and combine with the concept or
utility of the invention. Define any terms that may have a special meaning. Give examples or relevant citations, authors, etc, if
known. Please attach a copy of the cover sheet, pertinent claims, and abstract.
Title of Invention: Use of biologically active Vit D Compos
Inventors (please provide full names):
1. /2//1999 (US Pet 6,35-8,939)
Earliest Priority Filing Date: Hayes Colleen et al.
*For Sequence Searches Only* Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the
appropriate serial number. Jan Delaval Reference Librarian Symplomes
jan.delaval@usptggov / the wethost of
Please Search for the method (IBO). Such
for inflammatory bowl disesse (IBO), Such
Q 1 CC 2
1. tania D. 1406-16-2) V
by using
(32222-66-3)
1d hydroxy Vit. 03. (Cl 21)
2) by using 1 d hydroxy Vit. D3. (cl 21) v
31) 130447-37-9
(19-nor) 1. d., 25 dilydroxy (cl 37)
4) is Called 19-nor for
Please set attached sheet.
You way elected
banks in your sent of Medline the
STAFF USE ONLY Type of Search Vendors and cost where applicable
Searcher: NA Sequence (#) STN
Searcher Phone #:
Searcher Location: Structure (#) Questel/Orbit
Date Searcher Picked Up: SIY (37 Bibliographic Dr.Link
Date Completed: 5 1 4: 100 Litigation Lexis/Nexis
Searcher Prep & Review Time: Fulltext Sequence Systems
Clerical Prep Time: Patent Family WWW/Internet
Online Time: Other Other Other
PTO-1590 (8-01)

=> fil reg FILE 'REGISTRY' ENTERED AT 15:44:58 ON .14 SEP 2002 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2002 American Chemical Society (ACS)

STRUCTURE FILE UPDATES: 13 SEP 2002 HIGHEST RN 450944-74-8 DICTIONARY FILE UPDATES: 13 SEP 2002 HIGHEST RN 450944-74-8

TSCA INFORMATION NOW CURRENT THROUGH MAY 20, 2002

Please note that search-term pricing does apply when conducting  ${\tt SmartSELECT}$  searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Calculated physical property data is now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

=> d ide can tot 148

L48 ANSWER 1 OF 4 REGISTRY COPYRIGHT 2002 ACS

RN 131918-61-1 REGISTRY

CN 19-Nor-9,10-secoergosta-5,7,22-triene-1,3,25-triol, (1.alpha.,3.beta.,7E,22E)- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN Paricalcitol

CN Zemplar

FS STEREOSEARCH

MF C27 H44 O3

SR CA

LC STN Files: ADISINSIGHT, ADISNEWS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, DDFU, DIOGENES, DRUGNL, DRUGPAT, DRUGU, DRUGUPDATES, EMBASE, IPA, MEDLINE, MRCK\*, PHAR, PROMT, SYNTHLINE, TOXCENTER, USAN, USPATFULL (\*File contains numerically searchable property data)

Absolute stereochemistry. Double bond geometry as shown.

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

51 REFERENCES IN FILE CA (1967 TO DATE)

51 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 137:109489

REFERENCE 2: 137:56786

Jan Delaval
Reference Librarian
šiotechnology & Chemical Library
CM1 1E07 – 703-308-4498
ian delaval@uspto.gov

ADISINSIGHT, ADISNEWS, BEILSTEIN\*, BIOSIS, BIOTECHNO, CA,

CANCERLIT, CAPLUS, CASREACT, CBNB, CHEMCATS, DDFU, DIOGENES, DRUGNL, DRUGPAT, DRUGU, DRUGUPDATES, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, PHAR, RTECS\*, SYNTHLINE, TOXCENTER, USAN, USPATFULL

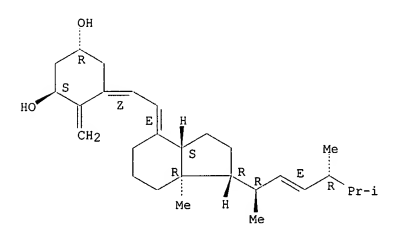
(\*File contains numerically searchable property data)

3: 136:355482 REFERENCE 136:345799 REFERENCE 4: 5: 136:319458 REFERENCE REFERENCE 6: 136:304095 REFERENCE 7: 136:273570 REFERENCE 8: 136:273235 136:241732 REFERENCE 9: REFERENCE 10: 136:161660 ANSWER 2 OF 4 REGISTRY COPYRIGHT 2002 ACS L48 RN 54573-75-0 REGISTRY 9,10-Secoergosta-5,7,10(19),22-tetraene-1,3-diol, CN (1.alpha., 3.beta., 5Z, 7E, 22E) - (9CI) (CA INDEX NAME) OTHER NAMES: CN 1-Hydroxyergocalciferol CN 1-Hydroxyvitamin D2 CN 1.alpha.-Hydroxyergocalciferol CN 1.alpha.-Hydroxyvitamin D2 CN Doxercalciferol CN Hectorol CN TSA 840 FS STEREOSEARCH 125285-48-5, 87649-67-0 DR MF C28 H44 O2

Absolute stereochemistry. Double bond geometry as shown.

STN Files:

LC



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

109 REFERENCES IN FILE CA (1967 TO DATE)

# 109 REFERENCES IN FILE CAPLUS (1967 TO DATE)

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REFERENCE
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REFERENCE
            2:
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            3:
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            4:
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REFERENCE
            5:
                136:289077
                136:273235
REFERENCE
            6:
REFERENCE
            7:
                136:241732
REFERENCE
            8:
                136:210551
REFERENCE
            9:
                136:161350
REFERENCE 10: 136:123683
L48 ANSWER 3 OF 4 REGISTRY COPYRIGHT 2002 ACS
RN
     41294-56-8 REGISTRY
     9,10-Secocholesta-5,7,10(19)-triene-1,3-diol, (1.alpha.,3.beta.,5Z,7E)-
CN
     (9CI) (CA INDEX NAME)
OTHER NAMES:
CN
     .alpha.-Calcidol
CN
     1-Hydroxycholecalciferol
CN
     1-Hydroxyvitamin D3
CN
     1.alpha.(OH)D3
CN
     1.alpha.-Hydroxycholecalciferol
CN
     1.alpha.-Hydroxyvitamin D3
CN
     Alfacalcidol
CN
     Alfarol
CN
     Alphacalcidol
CN
     Alpharol
CN
     Bondiol
CN
     Etalpha
CN
     Oxydevit
CN
     Un Alfa
CN
     Un Alpha
FS
     STEREOSEARCH
     125324-15-4, 41461-06-7, 43157-29-5, 43217-90-9
DR
MF
     C27 H44 O2
CI
     COM
LC
                  ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS,
       BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS,
       CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, DDFU, DRUGNL, DRUGPAT, DRUGU,
       DRUGUPDATES, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK*,
       NAPRALERT, PHAR, PHARMASEARCH, PROMT, RTECS*, SPECINFO, TOXCENTER, USAN,
       USPAT2, USPATFULL, VETU
         (*File contains numerically searchable property data)
     Other Sources:
                      EINECS**, WHO
         (**Enter CHEMLIST File for up-to-date regulatory information)
Absolute stereochemistry. Rotation (+).
Double bond geometry as shown.
```

HO 
$$\frac{Z}{Z}$$
 E  $\frac{H}{S}$   $\frac{R}{Me}$   $\frac{(CH_2)_3}{Me}$   $\frac{CHMe_2}{Me}$ 

# \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1068 REFERENCES IN FILE CA (1967 TO DATE)
23 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
1068 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 137:124547

REFERENCE 2: 137:120045

REFERENCE 3: 137:119679

REFERENCE 4: 137:119620

REFERENCE 5: 137:108703

REFERENCE 6: 137:59362

REFERENCE 7: 136:396366

REFERENCE 8: 136:345794

REFERENCE 9: 136:261073

REFERENCE 10: 136:230190

L48 ANSWER 4 OF 4 REGISTRY COPYRIGHT 2002 ACS

RN 1406-16-2 REGISTRY

CN Vitamin D (8CI, 9CI) (CA INDEX NAME)

MF Unspecified

CI COM, MAN

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMLIST, CIN, CSNB, DDFU, DIOGENES, DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, NAPRALERT, NIOSHTIC, PIRA, PROMT, RTECS\*, TOXCENTER, USPAT2, USPATFULL, VETU

(\*File contains numerically searchable property data)

Other Sources: EINECS\*\*, NDSL\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

#### \*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

6478 REFERENCES IN FILE CA (1967 TO DATE)

733 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

6487 REFERENCES IN FILE CAPLUS (1967 TO DATE)

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1: 137:174963
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            2:
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            4:
               137:168675
REFERENCE
            5:
               137:167659
REFERENCE
            6:
REFERENCE
            7:
               137:167593
               137:166985
REFERENCE
            8:
REFERENCE
            9:
               137:166979
REFERENCE 10: 137:166719
=> d his
     (FILE 'HOME' ENTERED AT 14:25:55 ON 14 SEP 2002)
                SET COST OFF
     FILE 'REGISTRY' ENTERED AT 14:26:21 ON 14 SEP 2002
                E VITAMIN D/CN
              1 S E3
L1
                STR
L2
             50 S L2 CSS
L3
     FILE 'HCAPLUS' ENTERED AT 14:28:01 ON 14 SEP 2002
                E HAYES C/AU
             39 S E3, E5
L4
                E HAYES COLEEN/AU
             52 S E4-E6
L5
                E NASHOLD F/AU
L6
             13 S E3-E6
            656 S (NORTH?(L)LIGHT?)/PA,CS
L7
            973 S (WISCON? (L) ALUM? (L) RES? (L) FOUND?) / PA, CS
^{18}
           6480 S L1
L9
          35010 S VITAMIN(S)D#
L10
L11
           5664 S ?CALCIFERO?
             14 S L4, L5, L6 AND L9-L11
L12
            159 S L7, L8 AND L9-L11
L13
              5 S L12 AND L13
L14
              9 S L12 NOT L14
L15
L16
           2619 S CALCITRIOL
           2418 S 1 ALPHA 25 DIHYDROXYVITAMIN D3
L17
           5759 S 1 25 DIHYDROXYVITAMIN D3
L18
             78 S 1 ALPHA 25 DIHYDROXYVITAMIN D2
L19
L20
             85 S 1 25 DIHYDROXYVITAMIN D2
              9 S 19 NOR 1 ALPHA 25 DIHYDROXYVITAMIN D2
L21
L22
              6 S 19 NOR 1 25 DIHYDROXYVITAMIN D2
             27 S PARICALCITOL
L23
     FILE 'REGISTRY' ENTERED AT 14:35:26 ON 14 SEP 2002
              3 S 32222-06-3 OR 60133-18-8 OR 131918-61-1
L24
     FILE 'HCAPLUS' ENTERED AT 14:38:03 ON 14 SEP 2002
L25
           9086 S L24
```

```
58 S ERCALCITRIOL OR ZEMPLAR OR RO176218 OR RO 17 6218 OR ROCALTRO
           1399 S (1 25 OR 1 ALPHA 25)()(DIHYDROXYCALCIFEROL OR DIHYDROXYERGOCA
L27
           3791 S (1 25 OR 1 ALPHA 25)()OH 2D3
L28
             68 S (1 25 OR 1 ALPHA 25)()OH 2D2
L29
          30838 S ?VITAMIN? ()(D OR D2 OR D3)
L30
          36555 S ?VITAMIN? (S) (D OR D2 OR D3)
L31
L32
          42102 S L10, L11, L16-L23, L26-31
          42200 S L32, L9, L25
L33
     FILE 'REGISTRY' ENTERED AT 14:42:36 ON 14 SEP 2002
              9 S (32222-06-3 OR 60133-18-8 OR 131918-61-1)/CRN
L34
     FILE 'HCAPLUS' ENTERED AT 14:43:10 ON 14 SEP 2002
             14 S L5-L6 AND L33
L35
                SEL RN
     FILE 'REGISTRY' ENTERED AT 14:44:03 ON 14 SEP 2002
L36
             23 S E1-E23
              3 S L36 AND L1, L24
L37
             20 S L36 NOT L37
L38
             18 S L38 AND C5-C6/ES AND C6/ES
L39
                SEL RN 12 18 17
L40
              3 S E24-E26
             15 S L39 NOT L40
L41
                E 1.ALPHA., 25-DIHYDROXYVITAMIN D3/CN
              1 S E3
L42
                E 19-NOR-1.ALPHA., 25-DIHYDROXYVITAMIN D2/CN
                E 1.ALPHA.-HYDROXYVITAMIN D3/CN
L43
              1 S E3
L44
              1 S E2
              3 S L1, L43, L44
L45
     FILE 'HCAPLUS' ENTERED AT 14:55:58 ON 14 SEP 2002
L46
             15 S L21, L22
     FILE 'REGISTRY' ENTERED AT 14:57:52 ON 14 SEP 2002
              1 S 131918-61-1
L47
              4 S L45, L47
L48
     FILE 'HCAPLUS' ENTERED AT 14:58:31 ON 14 SEP 2002
L49
           7460 S L48
             39 S PARICALCITOL OR ZEMPLAR OR L46
L50
             78 S DOXERCALCIFEROL OR HECTOROL OR TSA840 OR TSA 840 OR 1() (HYDRO
L51
            130 S ALPHA CALCIDOL OR ALFACALCIDOL OR ALFAROL OR ALPHACALCIDOL OR
L52
             36 S 1() (HYDROXYCHOLECALCIFEROL OR HYDROXYVITAMIN D3 OR OH D3)
L53
            .962 S 1()ALPHA() (HYDROXYCHOLECALCIFEROL OR HYDROXYVITAMIN D3 OR OH
L54
          20797 S VITAMIN D OR CALCIFEROL
L55
          21653 S L9, L49-L55
L56
             13 S L4-L6 AND L56
L57
                E INFLAMMATORY BOWEL/CT
                E E4+ALL
           2993 S E2
L58
                E INFLAMMATORY BOWEL/CT
                E E4+ALL
           3105 S INFLAMMATORY BOWEL() (DISEASE OR SYNDROME)
L59
L60
           1077 S IBD
                E ULCERATIVE COLITIS/CT
                E E3+ALL
           2115 S E2
L61
           3510 S ULCERATIVE ?COLITIS?
L62
                E CROHN/CT
                E E5+ALL
               0 S E2
L63
```

```
Page 7
```

```
1005 S CROHN?() (DISEASE OR ILEITIS OR INTESTIN? OR COLITIS)
 L64
 L65
              39 S L56 AND L58-L64
               1 S L57 AND L65
 L66
              23 S L65 AND (PY<=1999 OR PRY<=1999 OR AY<=1999)
 L67
              10 S (L49 OR L9) (L) (THU OR BAC OR USES) /RL AND L67
 L68
                 SEL DN AN 5 9
 L69
               2 S E1-E6
                 SEL DN AN L68 1-3
               3 S E7-E15
 L70
               '5 S L69,L70,L66 AND L4-L11,L16-L23,L25-L33,L35,L46,L49-L70
 L71
                 SEL RN L71 1
      FILE 'REGISTRY' ENTERED AT 15:37:03 ON 14 SEP 2002
 L72
              11 S E16-E26
 L73
               1 S L72 AND L48
              10 S L72 NOT L73
 L74
 L75
               9 S L74 NOT CA
      FILE 'HCAPLUS' ENTERED AT 15:37:45 ON 14 SEP 2002
                 E DIGESTIVE TRACT/CT
                 E E3+ALL
 L76
          141792 S E3, E101, E115
             320 S E66, E68, E69, E72
 L77
                 E COLITIS/CT
                 E E3+ALL
· L78
            3275 S E2
                 E INFLAMMATION/CT
            1308 S INFLAM?/CW (L) (INSTESTIN? OR BOWEL OR COLON? OR DIGEST? OR G
 L79
            2172 S L56 AND L76-L79
 L80
            2050 S L80 AND (PY<=1999 OR PRY<=1999 OR AY<=1999)
 L81
              39 S L81 AND (CROHN? OR ?ULCER? OR BOWEL OR COLIT?)
 L82
              19 S L82 NOT L65
 L83
               5 S L73, L75 AND L71
 L84
      FILE 'REGISTRY' ENTERED AT 15:44:58 ON 14 SEP 2002
 => d ide can tot 175
     ANSWER 1 OF 9 REGISTRY COPYRIGHT 2002 ACS
      346404-77-1 REGISTRY
 RN
      19-Nor-9, 10-secocholesta-5, 7, 15, 23-tetraene-1, 3, 25-triol,
 CN
      26,26,26-trifluoro-, (1.alpha.,7E,23E)- (9CI) (CA INDEX NAME)
 FS
      STEREOSEARCH
 MF
      C26 H37 F3 O3
 SR
      ÇA
                   CA, CAPLUS, USPATFULL
 LC
      STN Files:
 Absolute stereochemistry.
 Double bond geometry as shown.
```

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:56067

L75 ANSWER 2 OF 9 REGISTRY COPYRIGHT 2002 ACS

RN 153088-24-5 REGISTRY

CN 9,10-Secocholesta-5,7,10(19),16-tetraen-24-one, 1,3,25-trihydroxy-, (1.alpha.,2.beta.,5Z,7E)- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN JK 1624-3

CN Ro 25-8272

FS STEREOSEARCH

MF C27 H40 O4

SR CA

LC STN Files: CA, CAPLUS, CASREACT, TOXCENTER, USPATFULL

Absolute stereochemistry. Rotation (-). Double bond geometry as shown.

# \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

13 REFERENCES IN FILE CA (1967 TO DATE)

13 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:267379

REFERENCE 2: 135:116755

REFERENCE 3: 135:56067

REFERENCE 4: 135:28640

REFERENCE 5: 132:261062

REFERENCE 6: 132:30430

REFERENCE 7: 127:200524

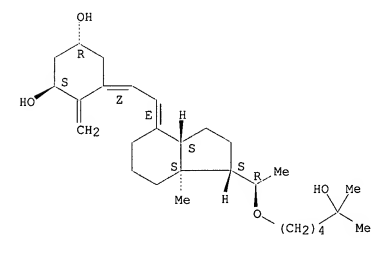
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REFERENCE 9: 126:195678

REFERENCE 10: 123:25141

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L75 ANSWER 3 OF 9 REGISTRY COPYRIGHT 2002 ACS
     132014-43-8 REGISTRY
RN
     1,3-Cyclohexanediol, 4-methylene-5-[(2E)-[(1S,3aS,7aS)-octahydro-1-[(1R)-1-1]]
CN
     [(5-hydroxy-5-methylhexyl)oxy]ethyl]-7a-methyl-4H-inden-4-
     ylidene]ethylidene]-, (1R, 3S, 5Z)- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     1,3-Cyclohexanediol, 4-methylene-5-[[octahydro-1-[1-[(5-hydroxy-5-
     methylhexyl)oxy]ethyl]-7a-methyl-4H-inden-4-ylidene]ethylidene]-,
     [1S-[1.alpha.(S*), 3a.beta., 4E(1S*, 3R*, 5Z), 7a.alpha.]]-
OTHER NAMES:
     KH 1049
CN
FS
     STEREOSEARCH
MF
     C28 H46 O4
SR
     CA
                  BEILSTEIN*, CA, CANCERLIT, CAPLUS, MEDLINE, TOXCENTER,
LC
     STN Files:
       USPATFULL
         (*File contains numerically searchable property data)
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Absolute stereochemistry. Double bond geometry as shown.



### \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

9 REFERENCES IN FILE CA (1967 TO DATE)

9 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:56067

REFERENCE 2: 130:20596

REFERENCE 3: 125:318815

REFERENCE 4: 122:283116

REFERENCE 5: 121:164013

REFERENCE 6: 120:125101

REFERENCE 7: 116:228269

REFERENCE 8: 116:75862

REFERENCE 9: 114:164629

L75 ANSWER 4 OF 9 REGISTRY COPYRIGHT 2002 ACS

RN 131875-08-6 REGISTRY

CN 1,3-Cyclohexanediol, 5-[(2E)-[(1S,3aS,7aS)-1-[(1R)-1-[(4-ethyl-4-hydroxyhexyl)oxy]ethyl]octahydro-7a-methyl-4H-inden-4-ylidene]ethylidene]-4-methylene-, (1R,3S,5Z)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 1,3-Cyclohexanediol, 5-[[1-[1-[(4-ethyl-4-hydroxyhexyl)oxy]ethyl]octahydro-7a-methyl-4H-inden-4-ylidene]ethylidene]-4-methylene-, [1S-[1.alpha.(S\*),3a.beta.,4E(1S\*,3R\*,5Z),7a.alpha.]]-

OTHER NAMES:

CN (5Z,7E,20R)-20-[(4-Ethyl-4-hydroxyhexyl)oxy]-9,10-secopregna-5,7,10(19)-triene-1.alpha.,3.beta.-diol

CN KH 106

CN KH 1060

CN Lexacalcitol

FS STEREOSEARCH

DR 138876-52-5

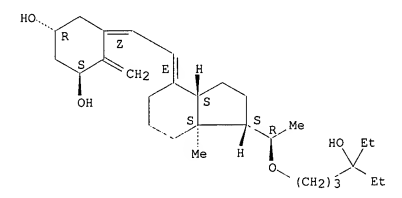
MF C29 H48 O4

SR CA

LC STN Files: ADISINSIGHT, ADISNEWS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, DDFU, DRUGNL, DRUGU, DRUGUPDATES, EMBASE, IPA, MEDLINE, PHAR, PROMT, SYNTHLINE, TOXCENTER, USAN, USPATFULL
Other Sources: WHO

Absolute stereochemistry.

Double bond geometry as shown.



## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

131 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

132 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 137:88763

REFERENCE 2: 137:33310

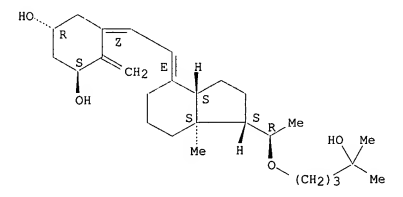
REFERENCE 3: 137:33309

REFERENCE 4: 136:379721

5: 136:241643 REFERENCE 136:211066 REFERENCE 6: 135:313933 REFERENCE 7: 135:298454 REFERENCE 135:236592 REFERENCE 9: REFERENCE 10: 135:205895 ANSWER 5 OF 9 REGISTRY COPYRIGHT 2002 ACS L75 131875-07-5 REGISTRY RN 1,3-Cyclohexanediol, 4-methylene-5-[(2E)-[(1S,3aS,7aS)-octahydro-1-[(1R)-1-1]]CN [(4-hydroxy-4-methylpentyl)oxy]ethyl]-7a-methyl-4H-inden-4ylidene]ethylidene]-, (1R, 3S, 5Z)- (9CI) (CA INDEX NAME) OTHER CA INDEX NAMES: 1,3-Cyclohexanediol, 4-methylene-5-[[octahydro-1-[1-[(4-hydroxy-4methylpentyl)oxy]ethyl]-7a-methyl-4H-inden-4-ylidene]ethylidene]-, [1S-[1.alpha.(S\*),3a.beta.,4E(1S\*,3R\*,5Z),7a.alpha.]]-OTHER NAMES: CN KH 1059 FS STEREOSEARCH C27 H44 O4 MF CA SR BEILSTEIN\*, BIOSIS, CA, CAPLUS, TOXCENTER, USPATFULL LC STN Files:

Absolute stereochemistry.

Double bond geometry as shown.



# \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

10 REFERENCES IN FILE CA (1967 TO DATE)
10 REFERENCES IN FILE CAPLUS (1967 TO DATE)

(\*File contains numerically searchable property data)

REFERENCE 1: 135:56067
REFERENCE 2: 130:20596
REFERENCE 3: 128:279055
REFERENCE 4: 125:266142

REFERENCE 5: 125:265006

REFERENCE 6: 120:125101

REFERENCE 7: 119:63329

REFERENCE 8: 116:228269

REFERENCE 9: 116:75862

REFERENCE 10: 114:164629

L75 ANSWER 6 OF 9 REGISTRY COPYRIGHT 2002 ACS

RN 124409-58-1 REGISTRY

CN 9,10-Secocholesta-5,7,10(19),16-tetraene-1,3,25-triol,

(1.alpha., 3.beta., 5Z, 7E) - (9CI) (CA INDEX NAME)

OTHER NAMES:

CN .DELTA.16-1.alpha., 25-Dihydroxyvitamin D3

CN 1.alpha., 25-Dihydroxy-. DELTA. 16-vitamin D3

CN 1.alpha., 25-Dihydroxy-16-ene-vitamin D3

CN 1.alpha., 25-Dihydroxyvitamin-16-ene D3

CN Ro 24-2637

CN VD 2708

FS STEREOSEARCH

DR 136198-25-9

MF C27 H42 O3

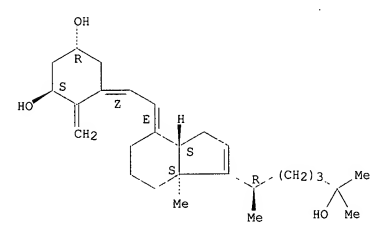
CI COM

SR CA

LC STN Files: CA, CANCERLIT, CAPLUS, MEDLINE, TOXCENTER, USPATFULL

Absolute stereochemistry.

Double bond geometry as shown.



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

50 REFERENCES IN FILE CA (1967 TO DATE)

2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

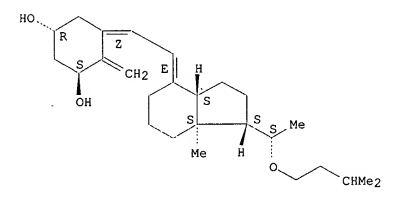
50 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:200351

REFERENCE 2: 135:116755

135:56484 REFERENCE 3: 135:56067 REFERENCE 4: 135:29446 REFERENCE 5: 135:28640 REFERENCE 6: 134:232195 REFERENCE 7: 134:217357 REFERENCE 8: 131:306845 REFERENCE 9: REFERENCE 10: 131:267036 L75 ANSWER 7 OF 9 REGISTRY COPYRIGHT 2002 ACS 111687-67-3 REGISTRY RN 1,3-Cyclohexanediol, 4-methylene-5-[(2E)-[(1S,3aS,7aS)-octahydro-7a-methyl-CN 1-[(1S)-1-(3-methylbutoxy)ethyl]-4H-inden-4-ylidene]ethylidene]-, (1R, 3S, 5Z) - (9CI)(CA INDEX NAME) OTHER CA INDEX NAMES: 1,3-Cyclohexanediol, 4-methylene-5-[2-[octahydro-7a-methyl-1-[1-(3methylbutoxy)ethyl]-4H-inden-4-ylidene]ethylidene]-, [1S-[1.alpha.(R\*), 3a.beta., 4E(1S\*, 3R\*, 5Z), 7a.alpha.]]-OTHER NAMES: CN 22-Oxa-1.alpha.-hydroxyvitamin D3 FS STEREOSEARCH DR 103909-46-2 MF C26 H42 O3 SR CA BEILSTEIN\*, CA, CANCERLIT, CAPLUS, CASREACT, MEDLINE, LC STN Files: TOXCENTER, USPATFULL (\*File contains numerically searchable property data)

Absolute stereochemistry.
Double bond geometry as shown.



# \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

9 REFERENCES IN FILE CA (1967 TO DATE)

9 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:56067

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130:20596
REFERENCE
            2:
                112:70796
REFERENCE
            3:
                111:17143
REFERENCE
                110:69959
REFERENCE
            5:
                110:29130
REFERENCE
            6:
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            7:
                110:13599
                108:88191
REFERENCE
            8:
REFERENCE
                108:6275
L75 ANSWER 8 OF 9 REGISTRY COPYRIGHT 2002 ACS
     57333-96-7 REGISTRY
RN
     9,10-Secocholesta-5,7,10(19)-triene-1,3,24-triol,
CN
     (1.alpha., 3.beta., 5Z, 7E, 24R) - (9CI) (CA INDEX NAME)
OTHER NAMES:
     1.alpha., 24(R) - Dihydroxycholecalciferol
CN
     1.alpha., 24(R)-Dihydroxyvitamin D3
CN
     1.alpha., 24R-Dihydroxyvitamin D3
CN
     Bonalfa
CN
     Curatoderm
CN
CN
     PRI 2191
CN
     Tacalcitol
CN
     TV 02
FS
     STEREOSEARCH
DR
     131801-95-1
     C27 H44 O3
MF
CI
                  ADISINSIGHT, ADISNEWS, AGRICOLA, BEILSTEIN*, BIOBUSINESS,
LC
       BIOSIS, CA, CAPLUS, CASREACT, CIN, DDFU, DRUGNL, DRUGPAT, DRUGU,
       DRUGUPDATES, IFICDB, IFIPAT, IFIUDB, MRCK*, PHAR, PROMT, RTECS*,
       SYNTHLINE, TOXCENTER, USAN, USPATFULL, VETU
         (*File contains numerically searchable property data)
     Other Sources:
```

Absolute stereochemistry. Double bond geometry as shown.

#### \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

176 REFERENCES IN FILE CA (1967 TO DATE)

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1 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
             176 REFERENCES IN FILE CAPLUS (1967 TO DATE)
            1: 137:114337
REFERENCE
               137:37660
REFERENCE
            2:
REFERENCE
            3:
                137:37408
                137:28324
REFERENCE
            4:
REFERENCE
            5:
                137:24157
REFERENCE
            6:
                137:24156
REFERENCE
            7:
                137:16058
                136:395928
REFERENCE
            8:
                136:345799
REFERENCE
            9:
REFERENCE 10: 136:226844
L75 ANSWER 9 OF 9 REGISTRY COPYRIGHT 2002 ACS
RN
     32222-06-3 REGISTRY
     9,10-Secocholesta-5,7,10(19)-triene-1,3,25-triol, (1.alpha.,3.beta.,5Z,7E)-
CN
      (9CI)
             (CA INDEX NAME)
OTHER NAMES:
     1,25-Dihydroxycholecalciferol
     1,25-Dihydroxyvitamin D
CN
CN
     1,25-Dihydroxyvitamin D3
     1.alpha., 25-(OH) 2D3
CN
     1.alpha., 25-Dihydroxycholecalciferol
CN
     1.alpha., 25-Dihydroxyvitamin D3
CN
CN
     Calcijex
CN
     Calcitriol
     Ro 21-5535
CN
CN
     Rocaltrol
CN
     Silkis
CN
     Soltriol
CN
     Topitriol
FS
     STEREOSEARCH
DR
     125338-24-1
     C27 H44 O3
MF
     COM
CI
                  ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN*,
LC
       BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT,
       CBNB, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU,
       DIOGENES, DRUGU, EMBASE, HSDB*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE,
       MRCK*, NAPRALERT, NIOSHTIC, PHAR, PHARMASEARCH, PROMT, RTECS*,
       TOXCENTER, USAN, USPAT2, USPATFULL, VETU
         (*File contains numerically searchable property data)
     Other Sources:
                      EINECS**, WHO
         (**Enter CHEMLIST File for up-to-date regulatory information)
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Absolute stereochemistry. Double bond geometry as shown.

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

9021 REFERENCES IN FILE CA (1967 TO DATE)
254 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
9031 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 137:174938

REFERENCE 2: 137:169690

REFERENCE 3: 137:169689

REFERENCE 4: 137:167659

REFERENCE 5: 137:167463

REFERENCE 6: 137:167289

REFERENCE 7: 137:166510

REFERENCE 8: 137:164102

REFERENCE 9: 137:164097

REFERENCE 10: 137:164062

### => fil hcaplus

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```
=> d all tot 184
    ANSWER 1 OF 5 HCAPLUS COPYRIGHT 2002 ACS
ΑN
    2001:472660 HCAPLUS
DN
    135:56067
ΤI
    Use of biologically active vitamin D compounds for the
    prevention and treatment of inflammatory bowel
IN
    Hayes, Colleen E.; Nashold, Faye E.
    Northern Lights Pharmaceuticals, LLC, USA
PA
SO
    PCT Int. Appl., 54 pp.
    CODEN: PIXXD2
DT
    Patent
LA
    English
    ICM C07C401-00
IC
    ICS A61K031-593
     1-7 (Pharmacology)
FAN.CNT 1
    PATENT NO.
                      KIND DATE
                                          APPLICATION NO. DATE
     _____
                      ____
                           _____
                                           -----
                            20010628
                                          WO 2000-US34913 20001221 <--
    WO 2001046132
                      A1
ΡI
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
             YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                           US 1999-469985
                       В1
                            20020319
                                                            19991221 <--
    US 6358939
    US 2002128241
                       A1
                            20020912
                                           US 2001-36819
                                                            20011221 <--
PRAI US 1999-469985
                            19991221
                                     <--
                       Α
OS
    MARPAT 135:56067
    Methods of treating inflammatory bowel disease
AB
    are described, and in particular the prevention and treatment of
    inflammatory bowel disease in humans as well
    as other animals. These methods involve the administration of biol.
    active vitamin D compds., and therapeutic compns.
    thereof, so that the symptoms of Inflammatory Bowel
    Disease are reduced or relieved.
ST
    vitamin D compd inflammatory bowel
    disease
IT
    Gene, animal
    RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (for inflammatory bowel disease risk;
       vitamin D compds. for prevention and treatment of
       inflammatory bowel disease)
IT
     Intestine, disease
        (inflammatory; vitamin D compds. for
        prevention and treatment of inflammatory bowel
        disease)
IT
    Drug delivery systems
```

```
(injections, i.v.; vitamin D compds. for prevention
        and treatment of inflammatory bowel disease
     Drug delivery systems
IΤ
        (oral; vitamin D compds. for prevention and
        treatment of inflammatory bowel disease)
ΙT
     Drug delivery systems
        (parenterals; vitamin D compds. for prevention and
        treatment of inflammatory bowel disease)
IT
     Drug delivery systems
        (rectal; vitamin D compds. for prevention and
        treatment of inflammatory bowel disease)
ΙT
     Drug delivery 'systems
        (topical; vitamin D compds. for prevention and
        treatment of inflammatory bowel disease)
ΙT
     Drug delivery systems
        (transdermal; vitamin D compds. for prevention and
        treatment of inflammatory bowel disease)
ΙT
     Intestine, disease
        (ulcerative colitis; vitamin D
        compds. for prevention and treatment of inflammatory
        bowel disease)
     Anti-inflammatory agents
    Cat (Felis catus)
     Dog (Canis familiaris)
     Horse (Equus caballus)
     Primate
        (vitamin D compds. for prevention and treatment of
        inflammatory bowel disease)
     Interleukin 10
ΙT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (vitamin D compds. for prevention and treatment of
        inflammatory bowel disease)
     1406-16-2, vitamin D 1406-16-2D,
    vitamin D, derivs. 32222-06-3,
     Calcitriol 57333-96-7 111687-67-3
     124409-58-1 131875-07-5 131875-08-6
     132014-43-8 153088-24-5 346404-77-1
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (vitamin D compds. for prevention and treatment of
        inflammatory bowel disease)
     7440-70-2, Calcium, biological studies
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (vitamin D compds. for prevention and treatment of
        inflammatory bowel disease)
              THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
(1) Calverley; US 5710142 A 1998 HCAPLUS
(2) Grue-Sorensen; US 5932565 A 1999 HCAPLUS
(3) Hesse; US 5786347 A 1998 HCAPLUS
(4) Schering Aktiengesellschaft; EP 0927721 Al 1999 HCAPLUS
    ANSWER 2 OF 5 HCAPLUS COPYRIGHT 2002 ACS
L84
     2001:435039 HCAPLUS
AN
DN
     135:41381
     Treatment of inflammatory bowel disease with
ΤI
     vitamin D compounds
     Cantorna, Margherita T.
IN
     The Penn State Research Foundation, USA
PA
```

```
SO
     PCT Int. Appl., 33 pp.
    CODEN: PIXXD2
DT
     Patent
LA
    English
IÇ
     ICM C07C401-00
CC
     2-10 (Mammalian Hormones)
FAN.CNT 1
                                           APPLICATION NO.
                                                            DATE
                      KIND DATE
     PATENT NO.
                                           _____
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                      ____
                                           WO 2000-US42393
                                                            20001130 <--
    WO 2001042205
                       A2
                            20010614
PΙ
     WO 2001042205
                       A3
                            20020321
             AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
             YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                           EP 2000-992552
                                                           20001130 <--
     EP 1233942
                       Α2
                            20020828
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
PRAI US 1999-168501P
                      Ρ
                            19991202
                                      <--
     US 2000-197827P
                       P
                            20000414
    US 2000-208632P
                       Ρ
                            20000601
    US 2000-231906P
                       Ρ
                            20000911
    WO 2000-US42393
                       W
                            20001130
OS
    MARPAT 135:41381
    A method of treating inflammatory bowel
AB
    disease, particularly ulcerative colitis and
     Crohn's disease, is disclosed. The method involves administering a
     vitamin D compd. in an amt. effective to treat the
     disease. The administration of a vitamin D compd.
     also prevents the development of or delays the onset of
     inflammatory bowel disease in susceptible
     individuals.
ST
     inflammatory bowel disease
    ulcerative colitis Crohns vitamin D
     treatment
IT
     Intestine, disease
        (Crohn's; treatment of inflammatory bowel
        disease with vitamin D compds.)
IT
     Intestine, disease
        (inflammatory; treatment of inflammatory
        bowel disease with vitamin D
        compds.)
IT
     Diet
        (low calcium; treatment of inflammatory bowel
        disease with vitamin D compds.)
ΙT
     Intestine, disease
        (ulcerative colitis; treatment of
        inflammatory bowel disease with
        vitamin D compds.)
     1406-16-2, vitamin D
IT
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (deficiency; treatment of inflammatory bowel
        disease with vitamin D compds.)
     32222-06-3D, analogs 41294-56-8, 1
ΙT
     .alpha.-Hydroxyvitamin D3 60133-18-8
     , 1,25-Dihydroxyvitamin D2
```

```
108646-38-4, 1.alpha.,25-
    75363-22-3
    Dihydroxyvitamin D3 triacetate 131918-61-1
    133876-00-3, 1.alpha.-Hydroxyvitamin D 156196-99-5
    195051-26-4
                  217093-03-3
    RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (treatment of inflammatory bowel disease
       with vitamin D compds.)
    7440-70-2, Calcium, biological studies
IT
    RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (treatment of inflammatory bowel disease
       with vitamin D compds.)
    ANSWER 3 OF 5 HCAPLUS COPYRIGHT 2002 ACS
    2001:255853 HCAPLUS
AN
DN
    134:271278
    Nutritional composition for treating inflammatory bowel
ΤI
    diseases
    Snowden, Robert B.
IN
    Snowden-Sutton Associates, Inc., USA
PA
SO
    U.S., 6 pp.
    CODEN: USXXAM
DT
    Patent
LA
    English
    ICM A61K047-00
IC
NCL 424439000
CC
     63-6 (Pharmaceuticals)
     Section cross-reference(s): 18
FAN.CNT 1
                                          APPLICATION NO. DATE
     PATENT NO.
                     KIND DATE
                      ____
                                          -----
                                                           19991007 <--
                                           US 1999-414666
ΡI
    US 6214373
                      В1
                            20010410
                                          WO 2000-US27404 20001005 <--
     WO 2001024642
                     A1
                           20010412
        W: CA
         RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE
PRAI US 1999-414666
                            19991007 <--
    A nutritional compn. and method useful for treatment of
    inflammatory bowel diseases is disclosed, the
     compn. comprising selected vitamins and mineral salts for oral
     administration to a subject having an inflammatory bowel
     disease. The compn. comprises an excess of vitamin
    D and vitamin B12 , contains vitamin C and
     iron in quantities promoting good absorption, contains water miscible
     forms of the fat-sol. vitamins, and no phosphate or carbonate
            Preferably, the iron is present as ferrous fumarate. And,
     preferably the compn. is essentially free of magnesium. Preferred compn.
     consists of retinyl acetate 2,500, cholecalciferol 400,
     dl-.alpha.-tocopherol acetate 75 IU, phytonadione 40 .mu.g, ascorbic acid
     100, thiamine mononitrate 5, riboflavin 5, pyridoxine hydrochloride 5 mg,
     cyanocobalamin 500 .mu.g, folic acid 0.2, niacinamide 10, biotin 0.15,
     pantothenic acid 5, iron 15, calcium 100, zinc 11.25 mg, selenium .mu.g,
     copper 1, manganese 1 mg, and iodine 75 .mu.g.
ST
     oral vitamin mineral inflammatory bowel
     disease
     Drug delivery systems
ΙT
        (caplets; vitamin and mineral compns. for treating inflammatory
        bowel diseases)
ΙT
     Drug delivery systems
        (capsules; vitamin and mineral compns. for treating
        inflammatory bowel diseases)
```

```
ΙT
     Intestine, disease
        (inflammatory; vitamin and mineral compns. for treating
        inflammatory bowel diseases)
IT
     Drug delivery systems
        (liqs., oral; vitamin and mineral compns. for treating
        inflammatory bowel diseases)
     Phosphates, biological studies
IT
     Sulfates, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (mineral; vitamin and mineral compns. for treating inflammatory
       bowel diseases)
IT
     Drug delivery systems
        (tablets; vitamin and mineral compns. for treating inflammatory
       bowel diseases)
IT
     Celiac disease
        (vitamin and mineral compns. for treating inflammatory
       bowel diseases)
IT
     Mineral elements, biological studies
     Vitamins
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (vitamin and mineral compns. for treating inflammatory
       bowel diseases)
IT
     9004-34-6, Cellulose, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (microcryst.; vitamin and mineral compns. for treating
        inflammatory bowel diseases)
     50-81-7, Ascorbic acid, biological studies
                                                  58-56-0, Pyridoxine
IT
     hydrochloride 58-85-5, Biotin 59-30-3, Folic acid, biological studies
                                              59-67-6, Niacin, biological
     59-43-8, Vitamin B1, biological studies
               67-97-0, Colecalciferol 68-19-9, Cyanocobalamin
     79-83-4, Pantothenic acid 83-88-5, Riboflavin, biological studies
     84-80-0, Phytonadione
                           98-92-0, Niacinamide 127-47-9, Retinyl acetate
     141-01-5, Ferrous fumarate
                                  532-43-4, Thiamine mononitrate
     1406-16-2, Vitamin D
                           1406-18-4, Vitamin E
     7439-89-6, Iron, biological studies
                                          7439-96-5, Manganese, biological
              7440-50-8, Copper, biological studies 7440-66-6, Zinc,
     biological studies 7440-70-2, Calcium, biological studies
     Iodine, biological studies 7782-49-2, Selenium, biological studies
     8059-24-3, Vitamin B6
                             9005-25-8, Starch, biological studies
                             12001-79-5, Vitamin K
                                                     52225-20-4, dl
     11103-57-4, Vitamin A
     -.alpha.-Tocopherol acetate
     RL: THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (vitamin and mineral compns. for treating
        inflammatory bowel diseases)
              THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
RE
(1) Anon; Gut 1986, V27(S1), P61
(2) Anon; Harrison's Principles of Internal Medicine 12th Ed 1991, V2, P1268
(3) Anon; Recommended Dietary Allowances 10th Ed 1989, P78
(4) Anon; Water-Soluble Vitamins P115
(5) Anon; Water-Soluble Vitamins P169
(6) Anon; Water-Soluble Vitamins P212
(7) Bennet; US 4617317 1986 HCAPLUS
(8) Brennan; US 4587793 1986
(9) Collins; The New England Journal of Medicine 1987, V316, P1654
(10) DeMichele; US 5780451 1998 HCAPLUS
(11) FernandezBanares; The American Journal of Gastroenterology 1989, V84(7),
    P744 MEDLINE
(12) Franklin; Impaired Folic Acid Absorption in Inflammatory Bowel Disease:
    Effects of Salicylazosulfapyridine 1973, V64(4), P517 MEDLINE
(13) Green; US 4806354 1989 HCAPLUS
```

(14) Harries; Postgraduate Medical Journal 1983, V59, P690 HCAPLUS

```
(15) Hathcock; Jama 1991, V265(1), P96 MEDLINE
(16) Hesse; US 5472957 1995 HCAPLUS
(17) Ivey; Handbook of Nonprescription Drugs 9th Ed 1990, P447
(18) Lashner; Gastroenterology 1989, V97, P255 MEDLINE
(19) Leddin; US 5578576 1996 HCAPLUS
(20) Lederle, F; Jama 1991, V265(1), P94 MEDLINE
(21) Linaker; Postgraduate Medical Journal 1979, V55, P26 MEDLINE
(22) McClain; Digestive Diseases and Sciences 1983, V28(1), P85 MEDLINE
(23) Nakamura; Digestive Diseases and Sciences 1988, V33(12), P1520 MEDLINE
(24) Nugent; American Gastroenterology Association 1979, V76(1), P1 MEDLINE
(25) Paradissis; US 5494678 1996 HCAPLUS
(26) Paul; US 5292538 1994 HCAPLUS
(27) Penny; Gut 1983, V24, P288 MEDLINE
(28) Peraita; US 5135918 1992 HCAPLUS
(29) Rosenberg; Gastroenterology 1989, V97, P502 MEDLINE
(30) Rowland; US 5405613 1995 HCAPLUS
(31) Sturniolo; Gut 1980, V21, P387 HCAPLUS
(32) Vogelsang; Digestive Diseases and Sciences 1989, V34(7), P1094 MEDLINE
    ANSWER 4 OF 5 HCAPLUS COPYRIGHT 2002 ACS
     2000:144772 HCAPLUS
ΑN
    132:189689
DN
ΤI
     Bioreductive conjugates for drug targeting
     Adams, Ged; Blake, David; Naughton, Declan; Stratford, Ian
IN
     Theramark Limited, UK; Adams, Margaret
PA
SO
     PCT Int. Appl., 48 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
     ICM A61K047-48
IC
CC
     1-12 (Pharmacology)
FAN.CNT 4
                  KIND DATE
     PATENT NO.
                                         APPLICATION NO. DATE
                           20000302 WO 1999-GB2606 19990819 <--
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                     A2
PΙ
    WO 2000010610
        W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,
            CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,
            IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD,
            MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK,
            SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY,
            KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
            ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
    AU 9954296
                     A1
                            20000314
                                          AU 1999-54296
                                                         19990819 <--
PRAI GB 1998-18027
                      Α
                            19980819
                            19980820
    GB 1998-18156
                      Α
                                     <--
                            19990819
    WO 1999-GB2606
                      W
                                     <--
    MARPAT 132:189689
OS
    The use of a bioreductive conjugate comprised of a noncytotoxic
AΒ
    bioreductive moiety having linked thereto at least one therapeutic agent,
     and salts thereof, is disclosed for the healing of wounds and the
     treatment of fibrotic disorders, ulcerative colitis,
     inflammatory bowel disease, epilepsy,
     cardiovascular reperfusion injury, cerebral reperfusion injury,
    hypertension, cystic fibrosis, psoriasis, para-psoriasis, peptic ulcers,
     gastric ulcers, duodenal ulcers, diabetic ulcers dementia, oncol., AIDS,
     rheumatoid arthritis, diabetes, and ischemia. Various specific conjugates
     for treating these conditions are also disclosed.
ST
    bioreductive conjugate drug targeting therapeutic
ΙT
     Transforming growth factors
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (TGF.beta.3; bioreductive conjugates for drug targeting)
```

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IT
     DNA
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (alkylation; bioreductive conjugates for drug targeting)
IT
     Psoriasis
        (and para-psoriasis; bioreductive conjugates for drug targeting)
IT
     Mitosis
        (antimitotics; bioreductive conjugates for drug targeting)
IT
     Actins
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (assembly and organization modulators; bioreductive conjugates for drug
        targeting)
ΙT
     Alkylation
        (biochem.; bioreductive conjugates for drug targeting)
IT
     Anti-AIDS agents
    Anti-inflammatory agents
    Anti-ischemic agents
    Anticoaqulants
    Anticonvulsants
     Antidiabetic agents
     Antihypertensives
     Antirheumatic agents
    Antitumor agents
    Antiulcer agents
    Apoptosis
    Cardiovascular agents
    Cystic fibrosis
     Drug metabolism
     Drug targeting
     Fibrinolytics
     Fibrosis
     Hypoxia, animal
     Immunomodulators
     Immunosuppressants
     Platelet aggregation inhibitors
     Radical scavengers
    Vasodilators
     Wound healing promoters
        (bioreductive conjugates for drug targeting)
TΤ
     Interleukin 10
     Interleukin 4
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (bioreductive conjugates for drug targeting)
TΤ
     Interleukin 1
     Platelet-derived growth factors
    Sex hormones
    RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (bioreductive conjugates for drug targeting)
IT
    Ion channel blockers
        (calcium; bioreductive conjugates for drug targeting)
ΙT
     Drugs
        (conjugates; bioreductive conjugates for drug targeting)
IT
    Corticosteroids, biological studies
    Steroids, biological studies
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (conjugates; bioreductive conjugates for drug targeting)
ΙT
     Diabetes mellitus
        (diabetic ulcer; bioreductive conjugates for drug targeting)
IT
    Cell cycle
```

```
(drugs specific for; bioreductive conjugates for drug targeting)
ΙT
     Intestine, disease
        (duodenum, ulcer; bioreductive conjugates for drug targeting)
IT
     Growth factors, animal
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (growth factor neutralizing agents; bioreductive conjugates for drug
        targeting)
IT
     Intestine, disease
        (inflammatory; bioreductive conjugates for drug targeting)
IT
     Lung, neoplasm
     Lung, neoplasm
        (inhibitors, A549; bioreductive conjugates for drug targeting)
TΤ
     Interleukin 6
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (inhibitors; bioreductive conjugates for drug targeting)
IT
     Reperfusion
        (injury, including cerebral reperfusion injury; bioreductive conjugates
        for drug targeting)
ΙT
     Integrins
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (integrin receptor activation inhibitors; bioreductive conjugates for
        drug targeting)
ΙT
     Antitumor agents
     Antitumor agents
        (lung, A549; bioreductive conjugates for drug targeting)
ΙT
     Ulcer
        (peptic; bioreductive conjugates for drug targeting)
IT
     Stomach, disease
        (ulcer; bioreductive conjugates for drug targeting)
IT
     Intestine, disease
        (ulcerative colitis; bioreductive conjugates for
        drug targeting)
IT
     Proteins, general, biological studies
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (wound site, growth factor-assocd.; bioreductive conjugates for drug
        targeting)
ΙT
     Adrenoceptor antagonists
        (.beta.-; bioreductive conjugates for drug targeting)
TT
     Polysaccharides, biological studies
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (.beta.-qlycans, sol.; bioreductive conjugates for drug targeting)
IT
     Transforming growth factors
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (.beta.1-; bioreductive conjugates for drug targeting)
ΙT
     Transforming growth factors
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (.beta.2-; bioreductive conjugates for drug targeting)
IT
     Interferons
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (.gamma.; bioreductive conjugates for drug targeting)
ΙT
     114560-25-7
                   114560-34-8, EO 8
                                       161518-24-7, RB 94547J
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); BIOL (Biological study)
        (bioreductive conjugates for drug targeting)
ΙT
     50-06-6D, Phenobarbitone, conjugates, biological studies
                                                                 50-24-8D,
     Prednisolone, conjugates
                                50-78-2D, Aspirin, conjugates
                                                                 52-53-9D,
     Verapamil, conjugates
                             52-67-5D, Penicillamine, conjugates
                                                                    53-86-1D,
     Indomethacin, conjugates
                                57-41-0D, Phenytoin, conjugates
                                                                   58-32-2D,
     Dipyridamole, conjugates
                                59-05-2D, Methotrexate, conjugates
                                                                      66-97-7D,
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89-57-6D, Mesalazine, conjugates

Psoralen, conjugates

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5-Aminosalicylic acid, derivs., conjugates 118-42-3D, Hydroxychloroquine, conjugates 305-03-3D, Chlorambucil, conjugates
    443-48-1D, Metronidazole, conjugates 446-86-6D, Azathioprine, conjugates
    599-79-1D, Sulfasalazine, conjugates
                                            1069-66-5D, Sodium valproate,
    conjugates 1406-16-2D, Vitamin D, analogs,
    conjugates 6556-11-2D, Inositol nicotinate, conjugates 12244-57-4D,
    Myochrysine, conjugates 15307-86-5D, Diclofenac, conjugates
    15687-27-1D, Ibuprofen, conjugates 21829-25-4D, Niphedipine, conjugates
    22204-53-1D, Naproxen, conjugates 26171-23-3D, Tolmetin, conjugates
    29679-58-1D, Fenoprofen, conjugates 38194-50-2D, Sulindac, conjugates
    51234-28-7D, Benoxaprofen, conjugates 56180-94-0D, Acarbose, conjugates 59865-13-3D, Cyclosporin A, conjugates 62571-86-2D, Captopril,
    conjugates 67763-97-7D, Insulin-like growth factor II, conjugates
     73590-58-6D, Omeprazole, conjugates 79217-60-0D, Cyclosporin, derivs.,
    conjugates 87333-19-5D, Ramipril, conjugates
                                                      87679-37-6D,
    Trandolapril, conjugates 97240-79-4D, Topiramate, conjugates
                                             113194-81-3, TMK 209
     103577-45-3D, Lansoprazole, conjugates
                                             259876-40-9, TMK 210
     117976-89-3D, Rabeprazole, conjugates
    259876-41-0, TMK 207
    RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (bioreductive conjugates for drug targeting)
     106096-92-8, Acidic fibroblast growth factor 106096-93-9, Basic
ΙT
     fibroblast growth factor
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (bioreductive conjugates for drug targeting)
     9015-82-1, Angiotensin-converting enzyme 9025-82-5, Phosphodiesterase
IT
     9036-21-9, Phosphodiesterase IV 9055-65-6, Prostaglandin synthetase
     9068-52-4, Phosphodiesterase V 81669-70-7, Metalloprotease 99676-46-7,
     Kexin 125978-95-2, Nitric oxide synthase
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (inhibitors; bioreductive conjugates for drug targeting)
     57285-09-3, Inhibin
                          114949-22-3, Activin
ΙT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (stimulators; bioreductive conjugates for drug targeting)
    ANSWER 5 OF 5 HCAPLUS COPYRIGHT 2002 ACS
L84
AN
    1998:668083 HCAPLUS
DN
     129:293874
     Pharmaceutical compositions containing flavonoids for the control and
ΤI
     treatment of anorectal and colonic diseases
     Singh, Amarjit; Jain, Rajesh; Singla, Anil Kumar
IN
     Panacea Biotec Ltd., India; University Institute of Pharmaceutical
PA
     Sciences
SO
     Eur. Pat. Appl., 17 pp.
     CODEN: EPXXDW
DT
     Patent
LA
     English
     ICM A61K031-35
IC
     ICS A61K031-70; A61K031-78
     63-5 (Pharmaceuticals)
     Section cross-reference(s): 1
FAN.CNT 1
                                           APPLICATION NO.
                                                             DATE
     PATENT NO.
                    KIND DATE
                     A1 19981007 EP 1997-302242 19970401 <--
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     EP 868914
PΙ
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
     A pharmaceutical compn., and process for the manuf. thereof, comprising
AB
     one or more flavonoids obtained from the plant Euphorbia prostata useful
     in the control and treatment of anorectal and colonic diseases.
     Standardized ext. of E. prostrata, when administered orally showed an
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inhibition of both carrageenan-induce edema with ED50 value of 5.98~mg/kg and histamine-induced edema with ED50 value of 16.37~mg/kg. A capsule contained above ext. 15, lactose 250, colloidal silicone dioxide 10, and talc 25 mg.

ST pharmaceutical capsule flavonoid anorectum colon disease

IT Balsams

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(Peru; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Quaternary ammonium compounds, biological studies

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(alkylbenzyldimethyl, chlorides; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Intestine

(anus, fissures; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Crack (fracture)

(anus; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Skin preparations (pharmaceutical)

(astringents; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Medical goods

(bandages; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(buccal; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(capsules; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Intestine, disease

Intestine, disease

(colon; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(films; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Digestive tract

(fistula; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(foams; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Vein

(hemorrhoid; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Intestine, disease

(inflammatory; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Irritants

(inhibitors; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(lozenges; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Keratins

RL: BSU (Biological study, unclassified); BIOL (Biological study) (lysis of, promoters of; pharmaceutical compns. contg. flavonoids for

qazi - 10 / 036819 control and treatment of anorectal and colonic diseases) Drug delivery systems TT (ointments, creams; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) IT Drug delivery systems (ointments; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) Drug delivery systems ΙT (pads; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) IT Drug delivery systems (parenterals; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) IT Abscess Anesthetics Antimicrobial agents Cholinergic antagonists Euphorbia prostrata Pigments, nonbiological Vasoconstrictors Wound healing Yeast (pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) IT Flavonoids RL: BAC (Biological activity or effector, except adverse); BOC (Biological occurrence); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); OCCU (Occurrence); USES (Uses) (pharmaceutical compns. contq. flavonoids for control and treatment of anorectal and colonic diseases) ΙT Castor oil Cocoa butter Cod liver oil Petrolatum Polyoxyalkylenes, biological studies Sterols Tannins Triterpenes RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) IT Alcohols, uses RL: NUU (Other use, unclassified); USES (Uses) (pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) ΙT Drug delivery systems (powders; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) IT Intestine (rectum, diseases; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) ΙT Fats and Glyceridic oils, biological studies RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (shark-liver oil; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases) Alkaloids, biological studies RL: BAC (Biological activity or effector, except adverse); BSU (Biological

study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES

(Uses)

(solanaceae; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(solns.; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(sprays; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(suppositories; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(suspensions; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(sustained-release; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Drug delivery systems

(tablets; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Intestine, disease

(ulcerative colitis; pharmaceutical compns. contq.

flavonoids for control and treatment of anorectal and colonic diseases)

IT Fats and Glyceridic oils, biological studies

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(vegetable; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT Hamamelis

(water; pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT 117-39-5, Quercetin 491-70-3, Luteolin 491-70-3D, Luteolin, glycoside derivs. 519-96-0D, 6-Methoxy quercetin, glycoside derivs. 520-36-5D, Apigenin, glycoside derivs.

RL: BAC (Biological activity or effector, except adverse); BOC (Biological occurrence); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); OCCU (Occurrence); USES (Uses)

(pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT 59-42-7, Phenylephrine 59-42-7D, Phenylephrine, salts 76-22-2, Camphor 85-79-0, Dibucaine 86-75-9, 8-Quinolinol benzoate 89-68-9, 89-78-1, Menthol 94-09-7, Benzocaine 94-24-6, Chlorothymol 99-26-3, Bismuth subgallate 101-08-6, Diperodon Tetracaine 101-93-9. Phenacaine 108-46-3, Resorcinol, biological studies 108-95-2, Phenol, 121-54-0, Benzethonium chloride biological studies 123-03-5, 134-31-6, 8-Hydroxyquinoline sulfate Cetylpyridinium chloride 299-42-3, Ephedrine 1314-13-2, Zinc oxide, 140-65-8, Pramoxine biological studies 1317-25-5 1406-16-2, Vitamin 8011-96-9, Calamine 8063-33-0 9005-25-8, Starch, 10043-35-3, Boric acid, biological studies biological studies

11103-57-4, Vitamin a 12263-41-1 21645-51-2, Aluminum hydroxide,

biological studies 25322-68-3, Peg 25322-69-4, Polypropylene glycol

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL

(Biological study); USES (Uses)

(pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

IT 141-78-6, Ethyl acetate, uses 7757-82-6, Sodium sulfate, uses RL: NUU (Other use, unclassified); USES (Uses)

(pharmaceutical compns. contg. flavonoids for control and treatment of anorectal and colonic diseases)

=> fil medline FILE 'MEDLINE' ENTERED AT 15:54:30 ON 14 SEP 2002

FILE LAST UPDATED: 13 SEP 2002 (20020913/UP). FILE COVERS 1958 TO DATE.

On June 9, 2002, MEDLINE was reloaded. See HELP RLOAD for details.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2002 vocabulary. Enter HELP THESAURUS for details.

THIS FILE CONTAINS CAS REGISTRY NUMBERS FOR EASY AND ACCURATE SUBSTANCE IDENTIFICATION.

=> d all tot

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L97 ANSWER 1 OF 6 MEDLINE
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AN 2001461932 MEDLINE

DN 20720280 PubMed ID: 11503842

TI Fractures in adults on systemic steroid therapy: which prophylaxis?.

AU Anonymous

SO Prescrire Int, (1999 Oct) 8 (43) 153-6. Journal code: 9439295. ISSN: 1167-7422.

CY France

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Health Technology

EM 200103

ED Entered STN: 20010820 Last Updated on STN: 20010820 Entered Medline: 20010329

(1) Systemic steroid therapy leads to a loss of bone density after a few AΒ months. The loss is at least partly reversible on treatment cessation. Together with age, the underlying disease, and reduced mobility, systemic steroid therapy is a risk factor for fractures. (2) There are no treatments with proven efficacy in the prevention of fractures among patients on systemic steroid therapy. Prevention is thus based on restricting steroid therapy to situations where the benefits are likely to outweigh the risks. (3) The first preventive measure is to encourage adequate calcium intake, as for all subjects at risk of osteoporosis. There is no firm evidence that all patients on steroids require medicinal calcium supplementation. (4) Some treatments slow the decline in bone density associated with steroid therapy, but none has a demonstrated preventive effect on symptomatic fractures. This is the case of the calcium + vitamin D combination, which has the best risk-benefit ratio. Two diphosphonates and, in postmenopausal women, hormone replacement therapy, also have a positive effect on bone density.

CT Check Tags: Comparative Study; Human Arthritis, Rheumatoid: DT, drug therapy

Asthma: DT, drug therapy

Bone Density

Calcitonin: TU, therapeutic use Calcium: TU, therapeutic use

Clinical Trials

Diphosphonates: TU, therapeutic use \*Fractures: CI, chemically induced

Inflammatory Bowel Diseases: DT, drug therapy

\*Osteoporosis: CI, chemically induced

Osteoporosis: DT, drug therapy

Osteoporosis: PC, prevention & control

\*Prednisone: AE, adverse effects Prednisone: TU, therapeutic use

Risk Factors

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Treatment Outcome
        Vitamin D: TU, therapeutic use
     1406-16-2 (Vitamin D); 53-03-2 (Prednisone); 7440-70-2
RN
     (Calcium); 9007-12-9 (Calcitonin)
     0 (Diphosphonates)
CN
     ANSWER 2 OF 6
L97
                       MEDLINE
                    MEDLINE
     1999215633
AN
               PubMed ID: 10201450
     99215633
DN
     Prevention and treatment of osteoporosis in patients with inflammatory
ΤI
     bowel disease.
ΑU
     Valentine J F; Sninsky C A
     Gainesville VA Medical Center and the Department of Medicine, University
CS
     of Florida 32610, USA.
     AMERICAN JOURNAL OF GASTROENTEROLOGY, (1999 Apr) 94 (4) 878-83.
SO
     Journal code: 0421030. ISSN: 0002-9270.
CY
     United States
     Journal; Article; (JOURNAL ARTICLE)
DT
     General Review; (REVIEW)
     (REVIEW, TUTORIAL)
     English
LA
FS
     Priority Journals
EΜ
     199904
     Entered STN: 19990511
ED
     Last Updated on STN: 19990511
     Entered Medline: 19990429
     Osteopenia or osteoporosis is common in patients with inflammatory bowel
AR
     disease. The use of corticosteroids contributes to the decline in bone
     loss; however, osteoporosis may develop in patients with inflammatory
     bowel disease independent of corticosteroid use. Risk factors for the
     development of low bone mass in patients with inflammatory bowel disease
     include the general risk factors for osteoporosis as well as additional
     factors such as the presence of chronic inflammation, use of
     corticosteroids and other pharmaceuticals, and nutritional deficiencies as
     the result of small bowel disease or small bowel resections. Despite the
     high prevalence, few patients are entered into prophylactic regimens to
     prevent corticosteroid-induced bone loss. The American College of
     Rheumatology has recently published recommendations for the prevention and
     treatment of corticosteroid-induced osteoporosis. In this article, we
     highlight the special risks for osteoporosis in patients with IBD and
     adapt the recommendations for prevention and treatment of osteoporosis to
     this clinical setting.
     Check Tags: Female; Human; Male
CT
      Anti-Inflammatory Agents, Steroidal: AE, adverse effects
      Bone Density
      Calcitonin: TU, therapeutic use
      Calcium Carbonate: TU, therapeutic use
      Diphosphonates: TU, therapeutic use
      Exercise
      Hormone Replacement Therapy
       *Inflammatory Bowel Diseases: CO, complications
        Inflammatory Bowel Diseases: EP, epidemiology
      Osteoporosis: EP, epidemiology
     *Osteoporosis: PC, prevention & control
      Prednisone: AE, adverse effects
      Risk Factors
        Vitamin D: TU, therapeutic use
     1406-16-2 (Vitamin D); 471-34-1 (Calcium Carbonate); 53-03-2
RN
     (Prednisone); 9007-12-9 (Calcitonin)
     0 (Anti-Inflammatory Agents, Steroidal); 0 (Diphosphonates)
CN
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L97 ANSWER 3 OF 6

MEDLINE

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AN
     1998415948
                    MEDLINE
                PubMed ID: 9744699
DN
     98415948
     A strategy for osteoporosis in gastroenterology.
ΤI
ΑU
     Scott E M; Scott B B
     Department of Endocrinology, St James's University Hospital, Leeds, UK.
CS
     EUROPEAN JOURNAL OF GASTROENTEROLOGY AND HEPATOLOGY, (1998 Aug)
SO
     10 (8) 689-96; discussion 696-8. Ref: 80
     Journal code: 9000874. ISSN: 0954-691X.
     ENGLAND: United Kingdom
CY
     Journal; Article; (JOURNAL ARTICLE)
DT
     General Review; (REVIEW)
     (REVIEW, TUTORIAL)
LA
     English
     Priority Journals
FS
     199811
EΜ
ED
     Entered STN: 19990106
     Last Updated on STN: 19990106
     Entered Medline: 19981119
     Osteoporotic fractures are a major public health problem.
AB
     Gastroenterologists see many patients at risk of osteoporosis,
     particularly those with coeliac disease and inflammatory bowel disease. In
     this paper, the extent of the problem is reviewed and a strategy of
     investigation and treatment is recommended.
     Check Tags: Female; Human; Male
CT
      Bone Density
      Densitometry, X-Ray
      Estrogen Replacement Therapy
      Fractures: ET, etiology
       *Inflammatory Bowel Diseases: CO, complications
      Mass Screening
     *Osteoporosis: CO, complications
      Osteoporosis: DI, diagnosis
     *Osteoporosis: PC, prevention & control
      Osteoporosis, Postmenopausal: PC, prevention & control
      Risk Factors
        Vitamin D: TU, therapeutic use
     1406-16-2 (Vitamin D)
RN
L97 ANSWER 4 OF 6
                       MEDLINE
                  MEDLINE
     96022523
ΑN
     96022523 PubMed ID: 8590154
DN
     Prevention of bone mineral loss in patients with Crohn's disease by
TI
     long-term oral vitamin D supplementation.
     Vogelsang H; Ferenci P; Resch H; Kiss A; Gangl A
ΠA
     Clinic of Internal Medicine IV (Department of Gastroenterology and
CS
     Hepatology), University of Vienna, Austria.
     EUROPEAN JOURNAL OF GASTROENTEROLOGY AND HEPATOLOGY, (1995 Jul)
SO
     7 (7) 609-14.
     Journal code: 9000874. ISSN: 0954-691X.
     ENGLAND: United Kingdom
CY
DΨ
     (CLINICAL TRIAL)
     Journal; Article; (JOURNAL ARTICLE)
     (RANDOMIZED CONTROLLED TRIAL)
LA
     English
FS
     Priority Journals
     199603
ĖΜ
     Entered STN: 19960404
ED
     Last Updated on STN: 19960404
     Entered Medline: 19960328
     OBJECTIVE: To determine whether long-term dietary supplementation with low
AB
     doses of vitamin D helps to prevent bone loss and the
     development of osteoporosis or osteomalacia in out-patients with Crohn's
     disease. DESIGN: A randomized controlled study. SETTING: The out-patient
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clinic of a tertiary centre (university hospital). PATIENTS: Seventy-five
out-patients (31 men and 44 women, aged 16-77 years) with Crohn's disease.
INTERVENTIONS: All patients were randomly assigned to receive either an
oral supplement of 1000 IU/day vitamin D for 1 year or
no supplement. Bone mineral density, assessed in the distal part of the
nondominant forearm using single photon absorptiometry, and serum levels
of 25-hydroxyvitamin D, assessed using a competitive protein binding
assay, were measured before and after the period of dietary
supplementation. MAIN OUTCOME MEASURE: Relative change of bone mineral
density. RESULTS: Serum levels of 25-hydroxyvitamin D increased in 57% of
patients who received a supplement (compared with 37% of control
patients). Bone mineral density decreased significantly in control
patients [median -7%, interquartile range -12.6-(+0.4%)] but not in
patients who received a supplement [median -0.2%, interquartile range
-3.8-(+14\%); P < 0.005]. Increases in bone mineral density were especially
prevalent among patients who received the supplement and had normal serum
levels of 25-hydroxyvitamin D (68%), whereas increases occurred in only
18% of patients with low serum levels of 25-hydroxyvitamin D (P = 0.008).
Patients without an intestinal resection and receiving the vitamin
D supplement had a marginally greater increase in bone mineral
content than patients who had undergone a resection (P = 0.05).
CONCLUSION: Long-term oral vitamin D supplementation
seems to be an efficient means of preventing bone loss in patients with
Crohn's disease and could be recommended, especially for patients at high
risk of osteoporosis.
Check Tags: Comparative Study; Female; Human; Male
 Absorptiometry, Photon
 Adult
 Bone Density
 Calcifediol: BL, blood
*Cholecalciferol: TU, therapeutic use
   Crohn Disease: CO, complications
  *Crohn Disease: DT, drug therapy
   Crohn Disease: ME, metabolism
 Osteomalacia: DI, diagnosis
*Osteomalacia: PC, prevention & control
 Osteoporosis: DI, diagnosis
*Osteoporosis: PC, prevention & control
 Time Factors
19356-17-3 (Calcifediol); 67-97-0 (Cholecalciferol)
ANSWER 5 OF 6
                  MEDLINE
85190106
            MEDLINE
           PubMed ID: 3991404
85190106
Symptomatic hypercalcaemia precipitated by magnesium therapy.
Nanji A A
POSTGRADUATE MEDICAL JOURNAL, (1985 Jan) 61 (711) 47-8.
Journal code: 0234135. ISSN: 0032-5473.
ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)
English
Priority Journals
198505
Entered STN: 19900320
Last Updated on STN: 19970203
Entered Medline: 19850528
A patient with Crohn's disease receiving vitamin D and
calcium had normal serum calcium levels when serum magnesium was low.
Hypercalcaemia was precipitated when supplemental magnesium was given. The
reason why serum calcium was initially normal is probably related to the
effect of magnesium deficiency in reducing serum calcium level.
Check Tags: Case Report; Female; Human
```

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Crohn Disease: DT, drug therapy
      Hypercalcemia: BL, blood
     *Hypercalcemia: CI, chemically induced
      Magnesium: BL, blood
     *Magnesium Sulfate: AE, adverse effects
        Vitamin D: TU, therapeutic use
     1406-16-2 (Vitamin D); 7439-95-4 (Magnesium); 7487-88-9
RN
     (Magnesium Sulfate)
    ANSWER 6 OF 6
                       MEDLINE
L97
     83028394
                  MEDLINE
ΑN
DN
     83028394
                PubMed ID: 6982188
    Vitamin D deficiency and bone disease in patients with
TI
     Crohn's disease.
     Driscoll R H Jr; Meredith S C; Sitrin M; Rosenberg I H
ΑU
SO
     GASTROENTEROLOGY, (1982 Dec) 83 (6) 1252-8.
     Journal code: 0374630. ISSN: 0016-5085.
CY
     United States
     Journal; Article; (JOURNAL ARTICLE)
DT
LA
    English
    Abridged Index Medicus Journals; Priority Journals
FS
EM
    198212
ED
     Entered STN: 19900317
    Last Updated on STN: 19900317
     Entered Medline: 19821218
     The prevalence of vitamin D deficiency in Crohn's
AB
     disease and the relationship of vitamin D status to
    metabolic bone disease have not been fully characterized. Serum
     25-hydroxyvitamin D was measured in 82 patients with Crohn's disease; 65%
     of Crohn's disease patients had a low serum 25-hydroxyvitamin D
     concentration; 25% had deficient levels (less than 10 ng/ml). The lowest
     25-hydroxyvitamin D levels were observed in patients with previous ileal
     resections. Nine patients were studied in detail including transiliac
    needle bone biopsies; 6 had osteomalacia and 3 osteoporosis. Six patients
    had repeat bone biopsies 9 to 18 mo after vitamin D
     treatment. Three patients with osteomalacia and low serum
     25-hydroxyvitamin D levels showed histologic improvement after therapy
    with oral vitamin D restored serum 25-hydroxyvitamin D
     levels to normal. The adequacy of therapy was assessed accurately by
    monitoring serum 25-hydroxyvitamin D concentration. Three patients with
    metabolic bone disease with normal serum 25-hydroxyvitamin D levels at
    diagnosis did not show histologic improvement after receiving
     vitamin D.
CT
    Check Tags: Female; Human; Male
      25-Hydroxyvitamin D 2
     Adult
     Aged
     *Bone Diseases, Metabolic: CO, complications
     Bone Diseases, Metabolic: DT, drug therapy
     Bone Diseases, Metabolic: PA, pathology
     Bone and Bones: PA, pathology
        Crohn Disease: BL, blood
       *Crohn Disease: CO, complications
        Crohn Disease: PA, pathology
      Ergocalciferols: AA, analogs & derivatives
     Ergocalciferols: BL, blood
     Middle Age
      Osteomalacia: CO, complications
        Vitamin D: TU, therapeutic use
       *Vitamin D Deficiency: CO, complications
        Vitamin D Deficiency: DT, drug therapy
     1406-16-2 (Vitamin D); 21343-40-8 (25-Hydroxyvitamin D 2)
RN
CN
     0 (Ergocalciferols)
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=> fil embase
FILE 'EMBASE' ENTERED AT 16:01:27 ON 14 SEP 2002
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 substance identification.
=> d all tot
L111 ANSWER 1 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.
     1999319084 EMBASE
ΤI
     [Demineralization of bone in Crohn's disease, its diagnosis, treatment and
     prevention).
     DEMINERALIZACE KOSTI U M. CROHN, JEJI DIAGNOSTIKA, LECBA A PREVENCE.
ΑU
     Kocian J.; Kocianova J.
CS
     Dr. J. Kocian, I Interni Klinika, IPVZ FTN, Videnska 800, 140 59 Praha 4,
     Czech Republic
SO
     Casopis Lekaru Ceskych, (1999) 138/17 (522-524).
     Refs: 30
     ISSN: 0008-7335 CODEN: CLCEAL
CY
     Czech Republic
DT
     Journal; General Review
FS
     033
             Orthopedic Surgery
     037
             Drug Literature Index
     038
             Adverse Reactions Titles
     048
             Gastroenterology
LA
     Czech
SL
     English; Czech
AB
     In 20 - 60% of patients with Crohn's disease bone demineralization is
     found, usually osteoporosis, but also osteoporosis with malatic features.
     The cause is the reduced calcium intake (loss of appetite, lactose
     intolerance and malabsorption), reduced vitamin D
     intake and corticoid therapy. Nowadays the diagnosis is facilitated by the
     use of densitometers (ultrasonic and DEXA) and markers of osteoresorption
     and new bone formation. In treatment in addition to calcium and
     vitamin D used for a long time, fluorides are
     administered (only as monofluorophosphate), nasal thyrocalcitonin and
     bisphosphonates of the third series (alendronate). In postmenopausal women
     also hormonal treatment can be used unless contraindicated. However,
     burdening of the bones with regular exercise is a necessity. For
     prevention adequate calcium and vitamin D intake is
     important, non-smoking, and exercise.
    Medical Descriptors:
       *Crohn disease: DT, drug therapy
     *osteoporosis: CO, complication
     *osteoporosis: DI, diagnosis
     *osteoporosis: DT, drug therapy
     *osteoporosis: PC, prevention
     *osteoporosis: SI, side effect
    bone demineralization: CO, complication
     bone demineralization: DT, drug therapy
    bone demineralization: PC, prevention
     bone demineralization: SI, side effect
     osteomalacia: CO, complication
     osteomalacia: DT, drug therapy
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osteomalacia: PC, prevention osteomalacia: SI, side effect

calcium intake corticosteroid therapy vitamin intake echography dual energy X ray absorptiometry hormonal therapy exercise human review Drug Descriptors: calcium: DT, drug therapy vitamin d: DT, drug therapy fluorophosphate: DT, drug therapy calcitonin: DT, drug therapy bisphosphonic acid derivative: DT, drug therapy alendronic acid: DT, drug therapy estrogen: DT, drug therapy gestagen: DT, drug therapy salcatonin: DT, drug therapy tridin: DT, drug therapy fluocalcic: DT, drug therapy corticosteroid: AE, adverse drug reaction corticosteroid: DT, drug therapy maxi kalz (calcium) 7440-70-2; (fluorophosphate) 10163-15-2, 15181-43-8, 7631-97-2, 7789-74-4; (calcitonin) 12321-44-7, 21215-62-3, 9007-12-9; (alendronic acid) 66376-36-1; (salcatonin) 47931-85-1 (1) Maxi kalz; (2) Fosamax; Fluocalcic; Miacalcic (1) Asta; (2) Merck Sharp and Dohme; Slovako; Biotika L111 ANSWER 2 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. 1998105778 EMBASE [Disorders of bone mineralization in Crohn's disease and their teatment]. PORUCHY MINERALIZACE KOSTI U CHROHNOVY CHOROBY A JEJICH LECBA. Kocian J.; Kocianova J. Dr. J. Kocian, I Interni Klinika, IPVZ FTN, Videnska 800, 140 59 Praha 4, Czech Republic Vnitrni Lekarstvi, (1998) 44/3 (162-165). Refs: 16 ISSN: 0042-773X CODEN: VNLEAH Czech Republic Journal; (Short Survey) 003 Endocrinology 037 Drug Literature Index Gastroenterology 048 Czech English; Czech Frequent complications of Crohn's disease include disorders of bone mineralization. They are due to a reduce dietary calcium supply in patients with lactose intolerance and a certain degree of malabsorption of calcium as well as vitamin D. The position is made worse by corticoids used in treatment of the basic disease, because they interfere not only with vitamin D conversation into its active (and much more effective) metabolites but also with osteoid formation. In the early diagnosis of demineralization a densitometer can be used; markers of bone metabolism are used so far less frequently. As to treatment either blockers of enhanced bone resorption can be used (Ca, vitamin D, bisposponates and thyrocalcitonin) or substances stimulating new formation of bone (F, growth factors, in postmenopause women hormonal substitution treatment) or a combination of preparations from both groups can be used. An irreplaceable part is played also by exercise, depending, of course, on the patient's general

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condition.

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CT
    Medical Descriptors:
       *crohn disease: DI, diagnosis
       *crohn disease: DT, drug therapy
     *osteoporosis: CO, complication
     *osteoporosis: DI, diagnosis
     *osteoporosis: DT, drug therapy
     *osteoporosis: EP, epidemiology
     *osteoporosis: TH, therapy
     lactose intolerance: DI, diagnosis
     lactose intolerance: DT, drug therapy
     malabsorption: ET, etiology
     densitometry
     hormone substitution
     exercise
     steroid therapy
     intranasal drug administration
     short survey
     Drug Descriptors:
     calcium: DT, drug therapy
       vitamin d: DT, drug therapy
     bisphosphonic acid derivative: DT, drug therapy
     calcitonin: DT, drug therapy
     growth factor: DT, drug therapy
     estrogen: CB, drug combination
     estrogen: DT, drug therapy
     gestagen: CB, drug combination
     gestagen: DT, drug therapy
     acetylsalicylic acid: DT, drug therapy
     salazosulfapyridine: DT, drug therapy
     dexamethasone: DT, drug therapy
     fluocalcic: DT, drug therapy
     tridin: DT, drug therapy
     alendronic acid: DT, drug therapy
     salcatonin: DT, drug therapy
     biomin h
     osteogenon
     (calcium) 7440-70-2; (calcitonin) 12321-44-7, 21215-62-3, 9007-12-9;
     (acetylsalicylic acid) 493-53-8, 50-78-2, 53663-74-4, 53664-49-6,
     63781-77-1; (salazosulfapyridine) 599-79-1; (dexamethasone) 50-02-2;
     (alendronic acid) 66376-36-1; (salcatonin) 47931-85-1
     Fosamax; Miacalcic; Biomin h; Osteogenon
CN
L111 ANSWER 3 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.
AN
     97261762 EMBASE
DN
     1997261762
    Medical therapy for inflammatory bowel disease.
TI
     Feagan B.G.; McDonald J.W.D.
AU
     Dr. B.G. Feagan, University of Western Ontario, Department of Medicine,
CS
     Division of Gastroenterology, London, Ont. N6A-5A5, Canada
     Current Opinion in Gastroenterology, (1997) 13/4 (307-311).
SO
     Refs: 33
     ISSN: 0267-1379 CODEN: COGAEK
CY
     United States
DТ
     Journal; (Short Survey)
             Health Policy, Economics and Management
FS
     036
     037
             Drug Literature Index
     048
             Gastroenterology
     English
LA
     English
SL
     In ulcerative colitis the results with a new preparation of budesonide
AB
     provide a model for development of topically active, orally administered
     compounds. This approach is promising for the treatment of intestinal
```

inflammation by this class of steroids, which are characterized by high potency and low systemic toxicity. Immunosuppressive treatment in ulcerative colitis remains a form of therapy whose role is uncertain pending large controlled studies that assess both efficacy and safety. For most patients with ulcerative colitis, 5-ASA remains a mainstay of chronic therapy. Although the use of newer mesalamine compounds is widely accepted among qastroenterologists, they appear to have only marginal benefits compared with sulphasalazine and are significantly more expensive. Economic analysis comparing these interventions is necessary. For Crohn's disease, oral steroid therapy remains the cornerstone of treatment and is substantially more effective than dietary therapy. The use of antibiotic therapy to induce remission requires further evaluation in large, randomized controlled trials. Immunosuppressive therapy with the purine antimetabolites or methotrexate is effective and safe for patients who are resistant to, or dependent on, steroid use. Medical Descriptors: \*enteritis: ET, etiology \*enteritis: DT, drug therapy

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Barri Y.M.; Graves G.S.; Knochel J.P.

\*enteritis: DM, disease management anemia: DR, drug resistance anemia: DT, drug therapy anemia: CO, complication antibiotic therapy crohn disease: DT, drug therapy crohn disease: TH, therapy diet therapy drug efficacy drug potency drug safety human immunosuppressive treatment nutrition osteopenia: CO, complication osteopenia: DT, drug therapy remission short survey steroid therapy ulcerative colitis: DT, drug therapy Drug Descriptors: antibiotic agent: DT, drug therapy budesonide: DT, drug therapy budesonide: PR, pharmaceutics erythropoietin: DT, drug therapy immunosuppressive agent: DT, drug therapy mesalazine: DT, drug therapy mesalazine: PE, pharmacoeconomics methotrexate: DT, drug therapy purine derivative: DT, drug therapy salazosulfapyridine: DT, drug therapy salazosulfapyridine: PE, pharmacoeconomics steroid: DT, drug therapy vitamin d: DT, drug therapy (budesonide) 51333-22-3; (erythropoietin) 11096-26-7; (mesalazine) 89-57-6; (methotrexate) 15475-56-6, 59-05-2, 7413-34-5; (salazosulfapyridine) 599-79-1 L111 ANSWER 4 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. AN 97149125 EMBASE 1997149125 Calciphylaxis in a patient with Crohn's disease in the absence of endstage renal disease.

Dr. J.P. Knochel, Department of Medicine, Presbyterian Hospital of Dallas,

```
8200 Walnut Hill Lane, Dallas, TX 75231, United States
     American Journal of Kidney Diseases, (1997) 29/5 (773-776).
SO
     Refs: 34
     ISSN: 0272-6386 CODEN: AJKDDP
CY
     United States
DT
     Journal; Article
FS
     003
             Endocrinology
     028
             Urology and Nephrology
     037
             Drug Literature Index
     English
LA
     English
SL
     Calciphylaxis is a rare and life-threatening condition of progressive
AB
     cutaneous necrosis secondary to small and medium-sized vessel
     calcification previously described in patients with end-stage renal
     disease and hyperparathyroidism. Early diagnosis may be important in
     improving the poor outcome in these patients since early intervention may
     forestall the development of life-threatening complications. We describe a
     patient with Crohn's disease complicated by short-bowel syndrome and
     modest renal insufficiency (not requiring renal replacement therapy) who
     developed calciphylaxis. It appears that longstanding Crohn's disease and
     the short- bowel syndrome accelerated the development of calciphylaxis as
     the chronic renal disease was not end stage. Considering the possibility
     of calciphylaxis in this setting may avoid delaying the diagnosis and its
     consequences.
CT
    Medical Descriptors:
     *calcinosis: CO, complication
     *calcinosis: PC, prevention
     *calcinosis: ET, etiology
     *calcinosis: DI, diagnosis
     *calcinosis: DT, drug therapy
     *chronic kidney failure: CO, complication
       *crohn disease: SU, surgery
       *crohn disease: DT, drug therapy
     *hyperphosphatemia: CO, complication
     *hyperphosphatemia: ET, etiology
     *hyperphosphatemia: DI, diagnosis
     *secondary hyperparathyroidism: SU, surgery
     *secondary hyperparathyroidism: CO, complication
     *secondary hyperparathyroidism: DI, diagnosis
     *secondary hyperparathyroidism: ET, etiology
     *short bowel syndrome: CO, complication
     adult
     article
     calcium blood level
     case report
     colon resection
     disease association
     early diagnosis
     female
     human
     parathyroid hormone blood level
     phosphate blood level
     postoperative complication
     Drug Descriptors:
     prednisone: DO, drug dose
     prednisone: DT, drug therapy
       vitamin d: DT, drug therapy
     (prednisone) 53-03-2
L111 ANSWER 5 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.
AN
     96124615 EMBASE
DN
     1996124615
TI
     Crohn's complicated by therapy.
```

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ΑU
     Parry T.
CS
     Aintree Hospitals, NHS Trust, Liverpool, United Kingdom
SO
     Pharmacy in Practice, (1996) 6/4 (131-132).
     ISSN: 0962-9734 CODEN: PHPRF7
CY
     United Kingdom
DT
     Journal; (Short Survey)
             Orthopedic Surgery
FS
     033
     048
             Gastroenterology
     030
             Pharmacology
     037
             Drug Literature Index
     038
             Adverse Reactions Titles
LA
     English
CT
     Medical Descriptors:
       *crohn disease: DI, diagnosis
       *crohn disease: DT, drug therapy
     *osteoporosis: SI, side effect
     *osteoporosis: CO, complication
     *osteoporosis: DT, drug therapy
     *osteoporosis: PC, prevention
     *osteoporosis: ET, etiology
     drug choice
     fracture
     human
     oral drug administration
     risk factor
     short survey
     symptomatology:
     Drug Descriptors:
     *corticosteroid: AE, adverse drug reaction
     *corticosteroid: DT, drug therapy
     *mesalazine: DT, drug therapy
     *salazosulfapyridine: DT, drug therapy
     alendronic acid: DT, drug therapy
     calcitonin: DT, drug therapy
     calcium salt: DT, drug therapy
     estrogen: DT, drug therapy
     etidronic acid: DT, drug therapy
       vitamin d: DT, drug therapy
     (mesalazine) 89-57-6; (salazosulfapyridine) 599-79-1; (alendronic acid)
     66376-36-1; (calcitonin) 12321-44-7, 21215-62-3, 9007-12-9; (etidronic
     acid) 2809-21-4, 3794-83-0, 58449-82-4, 7414-83-7
L111 ANSWER 6 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.
     95195315 EMBASE
AN
DN
     1995195315
     Osteoporosis, corticosteroids and inflammatory bowel disease.
ΤI
ΑU
     Compston J.E.
     Department of Medicine, Addenbrooke's Hospital, Cambridge CB2 2QQ, United
CS
     Kingdom
SO
     Alimentary Pharmacology and Therapeutics, (1995) 9/3 (237-250).
     ISSN: 0269-2813
                     CODEN: APTHEN
CY
     United Kingdom
DT
     Journal; General Review
FS
     003
             Endocrinology
     006
             Internal Medicine
     010
             Obstetrics and Gynecology
     020
             Gerontology and Geriatrics
     048
             Gastroenterology
     030
             Pharmacology
     037
             Drug Literature Index
     038
             Adverse Reactions TitlesDrug Literature Index
LA
     English
     English
SL
```

AB Osteoporosis is a serious complication of inflammatory bowel disease which has not received adequate recognition despite its high prevalence and potentially devastating clinical effects. Its pathogenesis remains poorly defined although corticosteroid therapy and sex hormone deficiency are likely to play a major role. Recent advances in the diagnosis and management of osteoporosis have facilitated early detection of bone loss and identified means by which this may be prevented. Bone density measurements to predict fracture risk and define thresholds for prevention and treatment should be performed routinely in patients with inflammatory disease. Hormone replacement therapy is effective in prevention of bone loss in peri- and post-menopausal patients, but the treatment of younger women and men of all ages requires further study. CT Medical Descriptors: \*enteritis: ET, etiology \*enteritis: DT, drug therapy \*hormone deficiency \*osteoporosis: DI, diagnosis \*osteoporosis: SI, side effect \*osteoporosis: DT, drug therapy \*osteoporosis: PC, prevention \*osteoporosis: ET, etiology bone density female hormone substitution human malnutrition menopause oral drug administration priority journal review ulcerative colitis: ET, etiology ulcerative colitis: DT, drug therapy vitamin deficiency Drug Descriptors: \*anabolic agent: DT, drug therapy \*bisphosphonic acid derivative: DT, drug therapy \*calcitonin: DT, drug therapy \*calcium: DT, drug therapy \*corticosteroid: AE, adverse drug reaction \*estrogen: DT, drug therapy \*fluoride sodium: DT, drug therapy \*gestagen: DT, drug therapy \*parathyroid hormone: DT, drug therapy \*vitamin d: DT, drug therapy etidronic acid: DT, drug therapy RN (calcitonin) 12321-44-7, 21215-62-3, 9007-12-9; (calcium) 7440-70-2; (fluoride sodium) 51668-54-3, 7681-49-4, 79933-27-0; (parathyroid hormone) 12584-96-2, 68893-82-3, 9002-64-6; (etidronic acid) 2809-21-4, 3794-83-0, 58449-82-4, 7414-83-7 L111 ANSWER 7 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. 95075839 EMBASE DN 1995075839 ΤI Decreased bone density in inflammatory bowel disease is related to corticosteroid use and not disease diagnosis. ΑU Bernstein C.N.; Seeger L.L.; Sayre J.W.; Anton P.A.; Artinian L.; Shanahan CS Section of Gastroenterology, Health Sciences Centre, University of Manitoba, 820 Sherbrook St, Winnipeg, Man. R3A-1R9, Canada SO Journal of Bone and Mineral Research, (1995) 10/2 (250-256). ISSN: 0884-0431 CODEN: JBMREJ

CY

DT

United States

Journal; Article

FS 031 Arthritis and Rheumatism Drug Literature Index 037 Adverse Reactions Titles 038 048 Gastroenterology English LA SLEnglish Although corticosteroid therapy is associated with the development of AΒ osteopenia, it is unclear whether the cause of osteopenia in inflammatory bowel disease (Crohn's disease and ulcerative colitis) is related to corticosteroid therapy or other disease-related variables. Patients with Crohn's disease (a diffuse gastrointestinal disease) could have greater osteopenia than patients with ulcerative colitis because of small bowel disease and secondary malabsorption of calcium and vitamin D. A cross- sectional analysis of consecutive patients with Crohn's disease and ulcerative colitis was undertaken. Bone density was determined by measurements of the L2-L4 spine, the total hip, and Ward's triangle using dual energy X-ray absorptiometry (DXA). A number of clinical parameters were recorded prior to bone density evaluation and analyzed by univariate and subsequently multivariate analysis to determine possible predictors of osteopenia. Of the 26 patients with Crohn's disease, diminished bone density (a Z score of at least -1) was found at the hip in 64% and at the spine in 44%; and of the 23 patients with ulcerative colitis diminished bone density was found at the hip in 43% and at the spine in 48%. Among all the variables tested, only corticosteroid use was a statistically significant predictor of diminished bone density (p = 0.025 for the spine and hip and p = 0.005 for Ward's triangle).Disease diagnosis (Crohn's disease compared with ulcerative colitis) did not predict or correlate with diminished bone density. No obvious associations were seen between the measurements of any serum hormones or biochemistries and bone density, although the patients using corticosteroids had lower serum calcium levels than the nonusers. Separate multivariate analyses were performed for males and females. Corticosteroid use was statistically significantly associated with diminished bone density in females but not in males. All patients with inflammatory bowel disease (both Crohn's disease and ulcerative colitis), independent of whether or not they have small bowel disease, who have been using corticosteroids for long periods should have their bone density status investigated, since they have a high prevalence of diminished bone density and, therefore, are at risk for bone fractures. Further studies are required to sort out factors that may make bone density in females more sensitive to the effects of corticosteroids than that of males. CTMedical Descriptors: \*bone density \*enteritis: DT, drug therapy \*enteritis: ET, etiology absorptiometry adult article calcium blood level clinical article controlled study crohn disease: DT, drug therapy crohn disease: ET, etiology female hormone determination human human cell human tissue male

Drug Descriptors:

osteopenia: SI, side effect

ulcerative colitis: ET, etiology ulcerative colitis: DT, drug therapy

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*aminosalicylic acid: DT, drug therapy
     *calcium: DT, drug therapy
     *corticosteroid: AE, adverse drug reaction
     *corticosteroid: DT, drug therapy
       *vitamin d: DT, drug therapy
     (aminosalicylic acid)^{-}133-10-8, 133-15-3, 28088-64-4, 51540-64-8, 65-49-6,
RN
     80702-32-5; (calcium) 7440-70-2
L111 ANSWER 8 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.
     95040561 EMBASE
ΑN
DN
     1995040561
     Metabolic bone assessment in patients with inflammatory bowel disease.
ΤI
     Abitbol V.; Roux C.; Chaussade S.; Guillemant S.; Kolta S.; Dougados M.;
ΑU
     Couturier D.; Amor B.
     Ctr. d'Evaluation Maladies Osseuses, Hopital Cochin, 27 rue du Faubourg
CS
     Saint Jacques, 75014 Paris, France
     Gastroenterology, (1995) 108/2 (417-422).
SO
     ISSN: 0016-5085 CODEN: GASTAB
CY
     United States
DT
     Journal; Article
             Drug Literature Index
FS
     037
     048
             Gastroenterology
LA
     English
SL
     English
     Background/Aims: Patients with inflammatory bowel disease are at risk for
AB
     osteopenia. To study the metabolic bone status of these patients, a cross-
     sectional study was conducted. Methods: Eighty-four patients (49 women, 35
     men) with inflammatory bowel disease, 34 of whom had Crohn's disease and
     50 ulcerative colitis (including 18 with prior coloproctectomy and
     ileoanal anastomosis), underwent clinical, dietary, and spine radiological
     assessments. Bone metabolism was assessed by measuring serum levels of
     calcium, phosphate, parathyroid hormone (1-84), 25-hydroxyvitamin D3,
     1,25- dihydroxyvitamin D3, and osteocalcin. Lumbar and femoral neck bone
     mineral densities were measured by dual energy X-ray absorptiometry.
     Results: Serum osteocalcin level was decreased in 29 patients (34%), 12 of
     whom had never undergone steroid therapy. The other biochemical markers of
     bone metabolism were in the normal range. Thirty-six patients (43%) had
     osteopenia, and 6 patients (7%) had vertebral crush fractures. Osteopenia
     was observed in 27 patients (52%) and 9 patients (28%) with and without
     corticosteroid therapy, respectively. No patient had clinical or
     biological signs of osteomalacia. Analysis of bone density (lumbar Z
     score) by a multiple regression analysis showed a statistically
     significant correlation with age, cumulative corticosteroid doses,
     sedimentation rate, and osteocalcin level (R2 = 0.76; P = 0.05).
     Conclusions: The results suggest that bone turnover in inflammatory bowel
     disease is characterized by low bone formation in the presence of normal
     levels of calcium-regulating hormones.
CT
     Medical Descriptors:
       *colon crohn disease: DT, drug therapy
       *colon crohn disease: SU, surgery
     *osteopenia: CO, complication
       *ulcerative colitis: SU, surgery
       *ulcerative colitis: DT, drug therapy
     adolescent
     adult
     aged
     article
     bone density
     bone mineralization
     bone turnover
     dose response
```

female human

```
ileoanal anastomosis
    major clinical study
    male
    ossification
    osteomalacia
    priority journal
    proctocolectomy
    vertebra fracture: CO, complication
    Drug Descriptors:
       *calcifediol: EC, endogenous compound
       *calcitriol: EC, endogenous compound
    *calcium ion: EC, endogenous compound
     *osteocalcin: EC, endogenous compound
     *parathyroid hormone: EC, endogenous compound
     *phosphate: EC, endogenous compound
    azathioprine: DO, drug dose
    azathioprine: DT, drug therapy
    mesalazine: DT, drug therapy
    salazosulfapyridine: DT, drug therapy
    steroid: DO, drug dose
    steroid: DT, drug therapy
       vitamin d: DT, drug therapy
     (calcifediol) 19356-17-3; (calcitriol) 32222-06-3, 32511-63-0, 66772-14-3;
RN
     (calcium ion) 14127-61-8; (osteocalcin) 136461-80-8; (parathyroid hormone)
     12584-96-2, 68893-82-3, 9002-64-6; (phosphate) 14066-19-4, 14265-44-2;
     (azathioprine) 446-86-6; (mesalazine) 89-57-6; (salazosulfapyridine)
     599-79-1
L111 ANSWER 9 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.
     91204437 EMBASE
AN
     1991204437
DN
     [Severe osteoporosis in a young woman with Crohn's disease].
TΤ
     SCHWERE OSTEOPOROSE BEI EINER JUNGEN PATIENTIN MIT MORBUS CROHN.
     Neef B.; Horing E.; Maier K.-E.; v. Gaisberg U.
AU
    Medizinische Klinik Bad Cannstatt, Priessnitzweg 24, W-7000 Stuttgart 50,
CS
     Germany
     Deutsche Medizinische Wochenschrift, (1991) 116/27 (1055-1060).
SO
     ISSN: 0012-0472 CODEN: DMWOAX
CY
     Germany
     Journal; Article
DT
             Internal Medicine
FS
     006
             Orthopedic Surgery
     033
     048
             Gastroenterology
             Drug Literature Index
     037
             Adverse Reactions Titles
     038
     German
LA
SL
     English
     Increasing pain in the region of the lumbar vertebrae occurred in a
AB
     23-year-old woman known for the past 61/2 years to have Crohn's disease
     affecting the ileocolon. Radiology revealed marked osteopenia with
     collapse and deformation of the vertebral bodies. The only pointer to a
     bone disease was a markedly lowered serum level of 25-OH-vitamin
     D (< 10 ng/ml). Biopsy from the ileal crest revealed pure
     osteoporosis without osteomalacia. Decisive pathogenetic factors were, in
     the main, glucocorticoid medication, malnutrition and the long duration of
     Crohn's disease. During treatment with monofluorophosphate, 152 g daily,
     in fixed combination with 600 mg calcium as well as calcitonin (initially
     100 I.U. daily subcutaneously for two weeks, than 100 I.U. every other day
     s.c.) and vitamin D (3 x 1,000 I.U. daily by mouth)
     she became free of symptoms, and she has remained so for 9 months.
     Medical Descriptors:
       *crohn disease: DT, drug therapy
     *osteoporosis: SI, side effect
```

```
*osteoporosis: CO, complication
     *osteoporosis: DT, drug therapy
       *vitamin d deficiency: CO, complication
     article
     bone biopsy
     case report
     female
     human
     lumbar spine
     malnutrition
     oral drug administration
     priority journal
     subcutaneous drug administration
     vitamin blood level
     adult
     Drug Descriptors:
       *25 hydroxyvitamin d: EC, endogenous compound
     *calcitonin: DT, drug therapy
     *calcitonin: CB, drug combination
     *calcium: DT, drug therapy
     *calcium: CB, drug combination
     *fluorophosphate: DT, drug therapy
     *fluorophosphate: CB, drug combination
     *glucocorticoid: AE, adverse drug reaction
       *vitamin d: DT, drug therapy
       *vitamin d: CB, drug combination
     (25 hydroxyvitamin d) 64719-49-9; (calcitonin) 12321-44-7, 21215-62-3,
RN
     9007-12-9; (calcium) 7440-70-2; (fluorophosphate) 10163-15-2, 15181-43-8,
     7631-97-2, 7789-74-4
L111 ANSWER 10 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.
    87090510 EMBASE
DN
     1987090510
     [Medical treatment of Crohn's disease].
ΤI
     TRAITEMENT MEDICAL DE LA MALADIE DE CROHN.
ΑU
     Service de Chirurgie Digestive, Hopital Saint-Antoine, 75012 Paris, France
CS
SO
     Gazette Medicale, (1987) 94/7 (41-47).
    CODEN: GAMEE8
CY
     France
DT
     Journal
FS
     037
             Drug Literature Index
LA
CT
     Medical Descriptors:
       *crohn disease
     *drug therapy
     therapy
     digestive system
     short survey
     human
     Drug Descriptors:
     *antibiotic agent
     *antiinflammatory agent
     *azathioprine
     *bcg vaccine
     *codeine
       *colecalciferol
     *colestyramine
     *cromoglycate disodium
     *cyanocobalamin
     *folic acid
     *levamisole
     *loperamide
```

```
*mesalazine
     *metronidazole
     *paregoric
     *prednisolone
     *prednisone
     *salazosulfapyridine
     (azathioprine) 446-86-6; (codeine) 76-57-3; (colecalciferol)
RN
     1406-16-2, 67-97-0; (colestyramine) 11041-12-6, 58391-37-0;
     (cromoglycate disodium) 15826-37-6, 16110-51-3, 93356-79-7, 93356-84-4;
     (cyanocobalamin) 53570-76-6, 68-19-9, 8064-09-3; (folic acid) 59-30-3,
     6484-89-5; (levamisole) 14769-73-4, 16595-80-5; (loperamide) 34552-83-5,
     53179-11-6; (mesalazine) 89-57-6; (metronidazole) 39322-38-8, 443-48-1;
     (paregoric) 8029-99-0; (prednisolone) 50-24-8; (prednisone) 53-03-2;
     (salazosulfapyridine) 599-79-1
CN
     Pentasa; Salazopyrin; Flagyl; Imurel; Questran
L111 ANSWER 11 OF 11 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.
     78036416 EMBASE
DN
     1978036416
ΤI
     [Drugs for treatment of gastrointestinal affections and for substitution
     therapy].
     ARZNEIMITTEL ZUR BEHANDLUNG VON MAGEN UND DARMERKRANKUNGEN SOWIE ZUR
     SUBSTITUTIONS THERAPIE.
ΑU
     Doelle W.
CS
     Med. Univ. Klin., Tubingen, Germany
SO
     Deutsche Apotheker Zeitung, (1977) 117/5 (164-165).
     CODEN: DAZEA2
DT
     Journal
FS
     037
             Drug Literature Index
LA
     German
CT
     Medical Descriptors:
     *clinical study
     *drug comparison
     *irritable colon
     *malabsorption
     *peptic ulcer
     *drug therapy
       *ulcerative colitis
     therapy
     major clinical study
     Drug Descriptors:
     *alpha tocopherol
     *antacid agent
     *antibiotic agent
     *calcium carbonate
     *carbenoxolone
     *carbonic acid
       *colecalciferol
     *colestyramine
     *cholinergic receptor blocking agent
     *cyanocobalamin
     *glucocorticoid
     *histamine receptor
     *iron
     *medium chain triacylglycerol
     *menadione
     *opiate
     *retinol
     *salazosulfapyridine
     *tranquilizer
RN
     (alpha tocopherol) 1406-18-4, 1406-70-8, 52225-20-4, 58-95-7, 59-02-9;
     (calcium\ carbonate)\ 13397-26-7,\ 13701-58-1,\ 14791-73-2,\ 471-34-1;
```

(carbenoxolone) 5697-56-3, 7421-40-1; (carbonic acid) 3812-32-6, 463-79-6;

CN

ΑN

DN

ΤI

ΑU CS

PΙ

SO

DT

LA

AΒ

CC

IT

IT

TΤ

IT

ΙT

RN

(colecalciferol) 1406-16-2, 67-97-0; (colestyramine) 11041-12-6, 58391-37-0; (cyanocobalamin) 53570-76-6, 68-19-9, 8064-09-3; (iron) 14093-02-8, 53858-86-9, 7439-89-6; (menadione) 58-27-5; (opiate) 53663-61-9, 8002-76-4, 8008-60-4; (retinol) 68-26-8, 82445-97-4; (salazosulfapyridine) 599-79-1 Biogastrone; Azulfidine => fil biosis FILE 'BIOSIS' ENTERED AT 16:06:53 ON 14 SEP 2002 COPYRIGHT (C) 2002 BIOLOGICAL ABSTRACTS INC. (R) FILE COVERS 1969 TO DATE. CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE. RECORDS LAST ADDED: 11 September 2002 (20020911/ED) => d all tot L123 ANSWER 1 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 2002:240461 BIOSIS PREV200200240461 Use of biologically active vitamin D compounds for the prevention and treatment of inflammatory bowel disease. Hayes, Colleen E. (1); Nashold, Faye E. (1) Madison, WI USA ASSIGNEE: Northern Lights Pharmaceuticals, LLC, Madison, WI, USA US 6358939 March 19, 2002 Official Gazette of the United States Patent and Trademark Office Patents, (Mar. 19, 2002) Vol. 1256, No. 3, pp. No Pagination. http://www.uspto.gov/web/menu/patdata.html. e-file. ISSN: 0098-1133. Patent English Methods of treating inflammatory bowel disease are described, and in particular the prevention and treatment of inflammatory bowel disease in humans as well as other animals. These methods involve the administration of biologically active vitamin D compounds, and therapeutic compositions thereof, so that the symptoms of Inflammatory Bowel Disease are reduced or relieved. NCL 514167000 Biochemical Studies - Sterols and Steroids \*10067 Pathology, General and Miscellaneous - Therapy \*12512 Digestive System - Physiology and Biochemistry \*14004 Digestive System - Pathology \*14006 Pharmacology - General \*22002 Pharmacology - Connective Tissue, Bone and Collagen - Acting Drugs \*22012 Pharmacology - Immunological Processes and Allergy \*22018 Major Concepts Pharmacology Parts, Structures, & Systems of Organisms bowel: digestive system Diseases inflammatory bowel disease: digestive system disease Chemicals & Biochemicals vitamin D: antiinflammatory - drug, biologically active, immunologic - drug Alternate Indexing Inflammatory Bowel Diseases (MeSH) 1406-16-2 (VITAMIN D)

```
L123 ANSWER 2 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
    1999:371250 BIOSIS
AN
    PREV199900371250
DN
    Osteoporosis as a risk in inflammatory bowel
ΤI
     Schoon, E.-J.; Wolffenbuttel, B. H. R.; Stockbrugger, R. W. (1)
ΑU
     (1) Dept. of Gastroenterology, Academic Hospital Maastricht, P. Debyelaan
CS
     25, 6202 AZ, Maastricht Netherlands
    Drugs of Today, (April, 1999) Vol. 35, No. SUPPL. A, pp. 17-28.
SO
    ISSN: 0025-7656.
DT
    General Review
LA
    English
    Digestive System - Pathology *14006
    Radiation - Radiation and Isotope Techniques *06504
    Biochemical Studies - Vitamins *10063
    Biochemical Studies - Sterols and Steroids *10067
    Biochemical Studies - Minerals *10069
    Anatomy and Histology, General and Comparative - Radiologic Anatomy
     *11106
     Pathology, General and Miscellaneous - Diagnostic *12504
    Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     Pharmacology - Connective Tissue, Bone and Collagen - Acting Drugs *22012
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508.
     Pathology, General and Miscellaneous - Therapy *12512
    Metabolism - Metabolic Disorders
    Nutrition - Minerals *13206
    Nutrition - Fat-Soluble Vitamins *13208
    Digestive System - General; Methods *14001
BC
    Hominidae
               86215
    Major Concepts
IT
        Gastroenterology (Human Medicine, Medical Sciences)
TT
     Diseases
        fracture: injury; inflammatory bowel
        disease: digestive system disease; metabolic bone disease: bone
        disease, metabolic disease; osteopenia: bone disease; osteoporosis:
        bone disease, diagnosis, treatment; Crohn's disease:
       digestive system disease, immune system disease
IT
     Chemicals & Biochemicals
       bisphosphonates: metabolic; calcium: supplementation; corticosteroids:
        antiinflammatory; vitamin D:
        supplementation
IT
    Alternate Indexing
       Bone Diseases, Metabolic (MeSH); Crohn Disease
        (MeSH); Fractures (MeSH); Inflammatory Bowel
        Diseases (MeSH); Osteoporosis (MeSH)
IT
    Methods & Equipment
        dual X-ray absorptiometry: diagnostic method
    Miscellaneous Descriptors
IT
       bone density
ORGN Super Taxa
       Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        human (Hominidae): patient
ORGN Organism Superterms
        Animals; Chordates; Humans; Mammals; Primates; Vertebrates
     7440-70-2 (CALCIUM)
RN
       1406-16-2 (VITAMIN D)
L123 ANSWER 3 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
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AN 1999:360592 BIOSIS

- qazi 10 / 036819 PREV199900360592 DN Osteoporosis in patients with inflammatory bowel TI disease - Prevalence and risk factors. Von Tirpitz, Ch. (1); Pischulti, G.; Klaus, J.; Rieber, A.; Brueckel, J.; ΑU Boehm, B. O.; Adler, G.; Reinshagen, M. (1) Abteilung Innere Medizin I, Universitaetsklinik Ulm, CS Robert-Koch-Strasse 8, D-89081, Ulm Germany Zeitschrift fuer Gastroenterologie, (Jan., 1999) Vol. 37, No. 1, SO pp. 5-12. ISSN: 0044-2771. DT Article LA German SL English; German Introduction: Osteopenia and osteoporosis are frequent but often AΒ underestimated complications in inflammatory bowel disease. In patients with IBD, several factors could contribute to osteopenia, but the pathogenetic mechanisms are still not completely understood. We carried out a prospective study to evaluate the prevalence and possible etiologic factors for osteopenia and subsequent osteoporosis in IBD-patients. Methods: 140 patients with inflammatory bowel disease (Crohn's disease n = 125, ulcerative colitis n = 15) underwent clinical and spine radiological assessments. Lumbar bone mineral densities were measured by dual energy X-ray absorptiometry (DXA). Markers of bone formation and resorption and vitamin D were assessed in n = 95 patients. Patients were asked about medication, previous or actual intestinal stenosis, smoking and intestinal resection. A lactose-H2-breath test was undertaken if lactose intolerance was clinically suspected. Results: Compared to age- and sex-matched healthy controls (Z-score), the prevalence of osteopenia (Z < -1) was 62%, while osteoporosis (Z < -2) occurred in 38%. The mean bone density of IBD-patients was osteopenic with no significant differences between Crohn's disease (Z = -1, 24) and ulcerative colitis (Z = -1, 25). Osteoporotic fractures were seen in three patients (2,1%). Crohn's disease patients with osteoporosis showed a significant lower body mass index (BMI) than patients with normal bone density. 52,9% of patients with manifest osteporosis underwent systemic steroid treatment in the preceeding year, but only 34% of those with normal bone density. Except hemoglobin, none of the biochemical markers showed a significant difference between osteoporosis, osteopenia and patients with normal bone density. Conclusion: The results show a high prevelance of osteopenia and osteporosis in IBD. Since
- still no standard treatment. The effect of osteoanabolic and antiresorptive agents must be evaluated in further studies.

  CC Digestive System Pathology \*14006
  Pathology, General and Miscellaneous Inflammation and Inflammatory Disease \*12508
  Bones, Joints, Fasciae, Connective and Adipose Tissue Physiology and Biochemistry \*18004
  Bones, Joints, Fasciae, Connective and Adipose Tissue Pathology \*18006
  Immunology and Immunochemistry General; Methods \*34502

osteoporosis is often associated with low body mass index, multiple

IBD is multifactorious and not completely understood, there is

intestinal resections and previous systemic steroid treatment, we suggest a bone densitometry in these patients. Since etiology of osteoporosis in

IT Major Concepts

Gastroenterology (Human Medicine, Medical Sciences); Orthopedics (Human Medicine, Medical Sciences)

IT Diseases

inflammatory bowel disease: bone
complications, digestive system disease; lactose intolerance:
congenital disease, metabolic disease, digestive system disease,
genetic disease; osteopenia: bone disease, etiology, risk factors,

prevalence; osteoporosis: bone disease, risk factors, etiology, prevalence; ulcerative colitis: bone complications, digestive system disease; Crohn's disease: bone complications, immune system disease, digestive system disease TΤ Alternate Indexing Bone Diseases, Metabolic (MeSH); Colitis, Ulcerative (MeSH); Crohn Disease (MeSH); Inflammatory Bowel Diseases (MeSH); Lactose Intolerance (MeSH); Osteoporosis (MeSH) ΙT Miscellaneous Descriptors body mass index; bone mineral density ORGN Super Taxa Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name human (Hominidae): patient ORGN Organism Superterms Animals; Chordates; Humans; Mammals; Primates; Vertebrates RN 63-42-3 (LACTOSE) L123 ANSWER 4 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 1998:74030 BIOSIS DN PREV199800074030 ΤI Inflammatory bowel disease and osteoporosis. Andreassen, H. (1); Rungby, J.; Dahlerup, J. F.; Mosekilde, L. ΑU CS (1) Dep. Internal Med., Roskilde County Hosp. Koge, DK-4600 Koge Denmark Scandinavian Journal of Gastroenterology, (Dec., 1997) Vol. 32, No. 12, pp. 1247-1255. ISSN: 0036-5521. DTArticle LA English The relation between inflammatory bowel disease (IBD) and osteoporosis has received increasing attention during the past decade. The prevalence of low bone mass in patients with EBD has been reported to be more than 50%. The development of a quick non-invasive method to diagnose osteoporosis (dual-energy X-ray absorptiometry) provides a practical tool to identify the patient who needs special attention. The aetiology of the bone disease in patients with IBD has still not been elucidated, but corticosteroids may play a major role. Studies on the prevention/treatment of IBD -related osteoporosis are scarce. In a single uncontrolled study hormone replacement therapy proved effective in preventing bone loss in peri- and post-menopausal women with IBD. A placebo-controlled study showed that supplementation with calcium and vitamin D prevents bone loss in patients with Crohn's disease. The present paper reviews our current knowledge on the mechanisms and epidemiology of IBD-related bone disease. Digestive System - Pathology \*14006 Biochemical Studies - Minerals \*10069 Pathology, General and Miscellaneous - Inflammation and Inflammatory Disease \*12508 Metabolism - Minerals \*13010 Metabolism - Metabolic Disorders \*13020 Nutrition - Water-Soluble Vitamins \*13210 Nutrition - Prophylactic and Therapeutic Diets \*13218 Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology \*18006 Pharmacology - Endocrine System \*22016 BC Hominidae 86215 IT Major Concepts Dental and Oral System (Ingestion and Assimilation); Skeletal System (Movement and Support) IT Diseases inflammatory bowel disease: digestive

system disease; osteoporosis: bone disease

Miscellaneous Descriptors low bone mass

ORGN Super Taxa

IT

Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia

ORGN Organism Name

human (Hominidae): patient

ORGN Organism Superterms

Animals; Chordates; Humans; Mammals; Primates; Vertebrates RN 7440-70-2 (CALCIUM)

1406-16-2 (VITAMIN D)

- L123 ANSWER 5 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 1996:532257 BIOSIS
- DN PREV199699254613
- TI A randomized, placebo-controlled trial of calcium supplementation for decreased bone density in corticosteroid-using patients with inflammatory bowel disease: A pilot study.
- AU Bernstein, C. N. (1); Seeger, L. L.; Anton, P. A.; Artinian, L.; Geffrey, S.; Goodman, W.; Belin, T. R.; Shanahan, F.
- CS (1) Sect. Gastroenterol., Univ. Manitoba, GB445 Health Science Cent., 820 Sherbrooke St., Winnipeg, MB R3A 1R9 Canada
- SO Alimentary Pharmacology & Therapeutics, (1996) Vol. 10, No. 5, pp. 777-786.
  ISSN: 0269-2813.
- DT Article
- LA English

AΒ

Background: Patients with inflammatory bowel disease (IBD) have a high prevalence of osteoporosis. A number of studies have found that corticosteroid use is associated with the development of osteoporosis in these patients. Calcium supplementation may be of benefit in corticosteroid-induced osteoporosis and calcium may be a nutrient that patients with IBD lack. Aim: To test the benefit of calcium supplementation on bone density in a pilot study over a 1-year period, in a group of corticosteroid-using patients with IBD, in a randomized, double-blind, placebo-controlled treatment study. Methods: Corticosteroid-using patients with IBD including males over the age of 18 years and premenopausal females, were randomized to receive either calcium carbonate 1000 mg plus vitamin D 250 IU (Oscal) or an identically matched placebo. Dual energy X-ray absorptiometry measurements of bone density were obtained at entry and at 1 year. At entry, and every 3 months thereafter, serum was collected for the measurement of haemoglobin, biochemistry and bone hormones. Simultaneously a 24-h urine collection was analysed for calcium excretion and creatinine clearance, and a 4-day food record was collected to document dietary calcium and vitamin D ingestion. Results: We found a high prevalence of moderately severe decreased bone density in corticosteroid-using patients with IBD. The dose of prednisone in the year prior to study entry was inversely correlated with bone density at the hip (R = -0.67, P = 0.004). At study entry serum osteocalcin was inversely correlated with corticosteroid dose in the year prior to the study (R = -0.64, P = 0.02) and at study end, directly correlated with the percentage change in spine bone density (R = 0.59, P =0.01). The dietary calcium intake of these patients was close to the current RDA (recommended daily intake) for premenopausal, post-adolescent adults. Calcium supplementation with small extra doses of vitamin D conferred no obvious benefit to bone density at the end of 1 year. There was no correlation between oral calcium ingestion and bone mass measurements. Both the treatment and placebo groups' bone density

remained relatively stable at 1 year, suggesting that bone loss in corticosteroid-using patients may peak early into the use of the corticosteroids. Conclusions: Calcium supplementation (1000 mg/day) conferred no significant benefit to bone density at 1 year in patients with corticosteroid-using IBD patients with osteoporosis. Future investigations should explore other therapeutic avenues that may have greater effects on increasing bone density in patients who already have considerable osteoporosis. Biochemical Studies - Sterols and Steroids 10067 Biochemical Studies - Minerals Pathology, General and Miscellaneous - Inflammation and Inflammatory Disease \*12508 Nutrition - Minerals \*13206 Digestive System - Pathology \*14006 Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology \*18006 Pharmacology - Digestive System \*22014 Pharmacology - Endocrine System \*22016 Toxicology - Pharmacological Toxicology \*22504 Hominidae \*86215 Major Concepts Gastroenterology (Human Medicine, Medical Sciences); Nutrition; Pathology; Pharmacology; Skeletal System (Movement and Support); Toxicology Chemicals & Biochemicals CALCIUM Miscellaneous Descriptors ADVERSE EFFECTS; ANTIINFLAMMATORY; BONE DISEASE; CALCIUM; CORTICOSTEROID; DECREASED BONE DENSITY; DIGESTIVE SYSTEM DISEASE; INFLAMMATORY BOWEL DISEASE; NO BENEFICIAL EFFECTS; NUTRITION; ORTHOPEDICS; OSTEOPOROSIS; PATIENT; PHARMACOLOGY; PILOT STUDY; RANDOMIZED, PLACEBO-CONTROLLED TRIAL; SUPPLEMENTATION; TOXICOLOGY ORGN Super Taxa Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name human (Hominidae) ORGN Organism Superterms animals; chordates; humans; mammals; primates; vertebrates 7440-70-2 (CALCIUM) L123 ANSWER 6 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 1996:249602 BIOSIS PREV199698805731 Asymptomatic malnutrition in children with inflammatory bowel disease. Kazlow, P.; Borger, C.; Cohn, L.; Collins, J.; Defelice, A.; Deckelbaum, R.; Narwal, S. Dep. Pediatr., Columbia Univ., New York, NY USA Pediatric Research, (1996) Vol. 39, No. 4 PART 2, pp. 120A. Meeting Info.: Joint Meeting of the American Pediatric Society and the Society for Pediatric Research Washington, D.C., USA May 6-10, 1996 ISSN: 0031-3998. Conference English General Biology - Symposia, Transactions and Proceedings of Conferences, Congresses, Review Annuals 00520 Biochemical Studies - Vitamins 10063 Biochemical Studies - Minerals 10069 Pathology, General and Miscellaneous - Inflammation and Inflammatory Disease \*12508 Nutrition - Malnutrition; Obesity \*13203 Nutrition - Minerals \*13206

BC IT

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Nutrition - Vitamins, General \*13207

Digestive System - Pathology \*14006 Pediatrics \*25000 Developmental Biology - Embryology - Morphogenesis, General \*25508 BC Hominidae \*86215 IT Major Concepts Development; Gastroenterology (Human Medicine, Medical Sciences); Nutrition; Pathology; Pediatrics (Human Medicine, Medical Sciences) IT Chemicals & Biochemicals VITAMIN A; VITAMIN D; VITAMIN E; CALCIUM ΙT Miscellaneous Descriptors CALCIUM; GROWTH FAILURE; MEETING ABSTRACT; NUTRIENT DEFICIENCY; VITAMIN A; VITAMIN D; VITAMIN E ORGN Super Taxa Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name human (Hominidae) ORGN Organism Superterms animals; chordates; humans; mammals; primates; vertebrates RN 68-26-8Q (VITAMIN A) 11103-57-40 (VITAMIN A) 1406-16-2 (VITAMIN D) 1406-18-4 (VITAMIN E) 7440-70-2 (CALCIUM) L123 ANSWER 7 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. AN 1996:120435 BIOSIS DN PREV199698692570 ΤI Relationships between vitamin D, parathyroid hormone and bone mineral density in inflammatory bowel disease. ΑU Silvennoinen, J. CS Gastroenterology Unit/Dep. Internal Med., Univ. Hosp. Oulu, SF-90220 Oulu SO Journal of Internal Medicine, (1996) Vol. 239, No. 2, pp. 131-137. ISSN: 0954-6820. DΨ Article LA English AB Objectives: To explore the relationships between vitamin D intake, serum parathyroid hormone (PTH) and 25-hydroxyvitamin D (250HD) concentrations, and bone mineral density (BMD) in inflammatory bowel disease (IBD). Setting: A university hospital clinic in Finland. Subjects: One hundred and fifty randomly selected patients with IBD from the hospital register and 73 healthy controls. Measurements: BMD of the lumbar spine and the proximal femur was measured with dual energy X-ray absorptiometry. Vitamin D intake and serum levels of 250HD and PTH were determined. Results: The IBD patients had a lower serum 25 OHD concentration (28.4 (SD 12.0) nmol L-1) than the controls (36.1 (16.7) nmol L-1; P = 0.001), whereas no differences in the vitamin D intake or the serum PTH levels were found. The serum 250HD concentrations and the vitamin D intake of the patients with ulcerative colitis (n = 67) were similar to those of the Crohn's disease patients (n = 76). The patients with Crohn's disease of the small bowel had slightly, but not significantly, lower serum 2 5 OHD concentrations (25.6 (11.0) nmol L-1) than the other Crohn's disease patients (31.4 (14.3) nmol L-1; P = 0.061). In the IBD patients, the vitamin D intake and the serum 25 OHD and PTH concentrations were not associated with BMD. Conclusions. Patients with IBD have lower serum levels of 25OHD than healthy controls, but similar serum PTH concentrations and vitamin D intake. Vitamin D intake. and the serum levels of 250HD and PTH are not associated with BMD, and malabsorption is unlikely to be a major factor in the aetiology of bone

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loss in unselected IBD patients.
 CC
      Biochemical Studies - Vitamins
                                        10063
      Biochemical Studies - Proteins, Peptides and Amino Acids
      Biochemical Studies - Minerals
                                       10069
      Pathology, General and Miscellaneous - Inflammation and Inflammatory
      Disease *12508
      Nutrition - Fat-Soluble Vitamins *13208
        Digestive System - Pathology *14006
      Endocrine System - Parathyroid *17010
      Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
 BC
      Hominidae *86215
 ΙT
      Major Concepts
         Endocrine System (Chemical Coordination and Homeostasis);
         Gastroenterology (Human Medicine, Medical Sciences); Nutrition;
         Pathology; Skeletal System (Movement and Support)
 IT
      Chemicals & Biochemicals
           VITAMIN D; PARATHYROID HORMONE
 IT
     Miscellaneous Descriptors
         MALABSORPTION; OSTEOPOROSIS
 ORGN Super Taxa
         Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
         human (Hominidae)
ORGN Organism Superterms
         animals; chordates; humans; mammals; primates; vertebrates
      1406-16-2 (VITAMIN D)
     9002-64-6 (PARATHYROID HORMONE)
L123 ANSWER 8 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1996:90355 BIOSIS
     PREV199698662490
     Osteoporosis in inflammatory bowel disease.
ΤĮ
ΑU
     Kraenzlin, M. E.
CS
     Missionsstr. 35, CH-4055 Basel Switzerland
SO
     Seibel, M. J. [Editor]; Kraenzlin, M. E. [Editor]. (1995) pp. 110-114.
     Osteoporosis. Osteoporose.
     Publisher: S. Karger AG P.O. Box, Allschwilerstrasse 10, CH-4009 Basel,
     Switzerland.
     Meeting Info.: First Interdisciplinary Osteoporosis Symposium Basel,
     Switzerland October 20-21, 1995
     ISBN: 3-8055-6248-9.
     Book; Conference
DT
LA
     German
CC
     General Biology - Symposia, Transactions and Proceedings of Conferences,
     Congresses, Review Annuals
                                  00520
     Biochemical Studies - Vitamins
                                       10063
     Biochemical Studies - Sterols and Steroids
                                                   10067
     Biochemical Studies - Minerals
                                      10069
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Metabolism - Minerals *13010
     Metabolism - Fat-Soluble Vitamins *13016
     Metabolism - Metabolic Disorders *13020
     Nutrition - Malnutrition; Obesity *13203
Nutrition - Fat-Soluble Vitamins *13208
       Digestive System - Pathology *14006
     Reproductive System - Physiology and Biochemistry *16504
     Endocrine System - Adrenals *17004
     Endocrine System - Gonads and Placenta * 17006
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
BC
     Hominidae *86215
IT
    Major Concepts
        Endocrine System (Chemical Coordination and Homeostasis);
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Gastroenterology (Human Medicine, Medical Sciences); Metabolism;
        Nutrition; Pathology; Reproductive System (Reproduction); Skeletal
        System (Movement and Support)
     Chemicals & Biochemicals
ΙT
        CALCIUM; VITAMIN D
ΙT
     Miscellaneous Descriptors
        BOOK CHAPTER; CALCIUM; ESTROGEN; GLUCOCORTICOID; MEETING PAPER;
        OSTEOMALACIA; PREVENTION; VITAMIN D
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        human (Hominidae)
ORGN Organism Superterms
        animals; chordates; humans; mammals; primates; vertebrates
     7440-70-2 (CALCIUM)
RN
       1406-16-2 (VITAMIN D)
L123 ANSWER 9 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1995:280681 BIOSIS
ΑN
DN
     PREV199598294981
ΤI
     Oral calcium and vitamin D does not impact on
     decreased bone density in inflammatory bowel
     disease (IBD): A prospective randomized
     placebo-controlled study.
     Bernstein, C. N. (1); Seeger, L. L.; Anton, P. A.; Artinian, L.; Goodman,
ΑU
    W. G.; Geffrey, S. P.; Belin, T.; Shanahan, F.
     (1) Dep. Med., Univ. Manitoba, Winnipeg, MB Canada
CS
    Gastroenterology, (1995) Vol. 108, No. 4 SUPPL., pp. A782.
SO
    Meeting Info.: 95th Annual Meeting of the American Gastroenterological
     Association and Digestive Disease Week San Diego, California, USA May
     14-17, 1995
     ISSN: 0016-5085.
DT
    Conference
LA
     English
    General Biology - Symposia, Transactions and Proceedings of Conferences,
CC
     Congresses, Review Annuals
     Biochemical Studies - Vitamins
     Biochemical Studies - Sterols and Steroids
                                                  10067
     Biochemical Studies - Minerals
                                      10069
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Pathology, General and Miscellaneous - Therapy
                                                      *12512
    Metabolism - Minerals *13010
     Nutrition - Minerals
                          *13206
     Nutrition - Prophylactic and Therapeutic Diets *13218
       Digestive System - Pathology *14006
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Pharmacology - Clinical Pharmacology
     Pharmacology - Endocrine System *22016
     Toxicology - Pharmacological Toxicology
                                               *22504
     Hominidae *86215
BC
IT
    Major Concepts
        Gastroenterology (Human Medicine, Medical Sciences); Metabolism;
        Nutrition; Pathology; Pharmacology; Skeletal System (Movement and
        Support); Toxicology
ΙT
    Chemicals & Biochemicals
        CALCIUM: VITAMIN D
IT
    Miscellaneous Descriptors
          ANTIINFLAMMATORY AGENT; CALCIUM SUPPLEMENTATION;
        CORTICOSTEROID; MEETING ABSTRACT
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
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human (Hominidae)
ORGN Organism Superterms
        animals; chordates; humans; mammals; primates; vertebrates
     7440-70-2 (CALCIUM)
RN
       1406-16-2 (VITAMIN D)
L123 ANSWER 10 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1995:275281 BIOSIS
AN
     PREV199598289581
DN
     Decreased bone density in inflammatory bowel
ΤI
     disease is related to corticosteroid use and not disease
     diagnosis.
     Bernstein, Charles N. (1); Seeger, Leanne L.; Sayre, James W.; Anton,
ΑU
     Peter A.; Artinian, Lucy; Shanahan, Fergus
     (1) Univ. Manitoba, Section Gastroenterology, Health Science Centre, Room
CS
     GG-449, 820 Sherbrook Street, Winnipeg, MB R3A 1R9 Canada
     Journal of Bone and Mineral Research, (1995) Vol. 10, No. 2, pp. 250-256.
SO
     ISSN: 0884-0431.
     Article
DT
     English
LA
     Although corticosteroid therapy is associated with the development of
AB
     osteopenia, it is unclear whether the cause of osteopenia in
     inflammatory bowel disease (Crohn's
     disease and ulcerative colitis) is related to
     corticosteroid therapy or other disease-related variables. Patients with
     Crohn's disease (a diffuse gastrointestinal disease) could have
     greater osteopenia than patients with ulcerative colitis
     because of small bowel disease and secondary malabsorption of calcium and
     vitamin D. A cross-sectional analysis of consecutive
     patients with Crohn's disease and ulcerative
     colitis was undertaken. Bone density was determined by
     measurements of the L2-L4 spine, the total hip, and Ward's triangle using
     dual energy X-ray absorptiometry (DXA). A number of clinical parameters
     were recorded prior to bone density evaluation and analyzed by univariate
     and subsequently multivariate analysis to determine possible predictors of
     osteopenia. Of the 26 patients with Crohn's disease, diminished
     bone density (a Z score of at least -1) was found at the hip in 64% and at
     the spine in 44%; and of the 23 patients with ulcerative
     colitis diminished bone density was found at the hip in 43% and at
     the spine in 48%. Among all the variables tested, only corticosteroid use
     was a statistically significant predictor of diminished bone density (p =
     0.025 for the spine and hip and p = 0.005 for Ward's triangle). Disease
     diagnosis (Crohn's disease compared with ulcerative
     colitis) did not predict or correlate with diminished bone
     density. No obvious associations were seen between the measurements of any
     serum hormones or biochemistries and bone density, although the patients
     using corticosteroids had lower serum calcium levels than the nonusers.
     Separate multivariate analyses were performed for males and females.
     Corticosteroid use was statistically significantly associated with
     diminished bone density in females but not in males. All patients with
      inflammatory bowel disease (both Crohn
      's disease and ulcerative colitis), independent of
     whether or not they have small bowel disease, who have been using
     corticosteroids for long periods should have their bone density status
      investigated, since they have a high prevalence of diminished bone density
      and, therefore, are at risk for bone fractures. Further studies are
      required to sort out factors that may make bone density in females more
      sensitive to the effects of corticosteroids than that of males.
      Biochemical Studies - Sterols and Steroids
CC
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Biochemical Studies - Minerals 10069
Pathology, General and Miscellaneous - Diagnostic \*12504
Pathology, General and Miscellaneous - Inflammation and Inflammatory
Disease \*12508

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Digestive System - Pathology *14006
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Pharmacology - Endocrine System *22016
     Toxicology - Pharmacological Toxicology
                                               *22504
     Hominidae *86215
BC
ΙT
     Major Concepts
        Gastroenterology (Human Medicine, Medical Sciences); Pathology;
        Pharmacology; Skeletal System (Movement and Support); Toxicology
ΙT
     Miscellaneous Descriptors
        BONE MINERAL DENSITY; CORTICOSTEROID TOXICITY; CROHN'S
        DISEASE; OSTEOPENIA; ULCERATIVE COLITIS
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        human (Hominidae)
ORGN Organism Superterms
        animals; chordates; humans; mammals; primates; vertebrates
L123 ANSWER 11 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1995:104895 BIOSIS
ΑN
     PREV199598119195
DN
     Vitamin and mineral supplementation in inflammatory
ТT
     bowel disease: Article eight in the series.
ΑIJ
     Mason, Joel B.
     Div. Clin. Nutrition, Tufts Univ. Sch. Med., Boston, MA USA
CS
     Practical Gastroenterology, (1994) Vol. 18, No. 11, pp. 18A-18D, 18F-18H.
SO
     ISSN: 0277-4208.
DT
     Article
LA
     English
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
CC
     Disease
               12508
     Pathology, General and Miscellaneous - Therapy
                                                       12512
     Metabolism - Minerals *13010
     Metabolism - Fat-Soluble Vitamins *13016
     Metabolism - Water-Soluble Vitamins *13018
     Nutrition - Minerals *13206
     Nutrition - Fat-Soluble Vitamins *13208
     Nutrition - Water-Soluble Vitamins *13210
       Digestive System - Pathology *14006
     Hominidae *86215
BC
TΤ
     Major Concepts
        Gastroenterology (Human Medicine, Medical Sciences); Metabolism;
        Nutrition
IT
     Chemicals & Biochemicals
        VITAMIN B-12; FOLATE; VITAMIN D; CALCIUM;
        MAGNESIUM; PHOSPHATE; ZINC; IRON
ΙT
     Miscellaneous Descriptors
        CALCIUM; FOLATE; IRON; MAGNESIUM; PHOSPHATE; VITAMIN B-12;
        VITAMIN D; ZINC
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        human (Hominidae)
ORGN Organism Superterms
        animals; chordates; humans; mammals; primates; vertebrates
RN
     68-19-9 (VITAMIN B-12)
     59-30-3 (FOLATE)
       1406-16-2 (VITAMIN D)
     7440-70-2 (CALCIUM)
     7439-95-4 (MAGNESIUM)
     14265-44-2 (PHOSPHATE)
     7440-66-6 (ZINC)
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7439-89-6 (IRON)

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L123 ANSWER 12 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1995:82716 BIOSIS
DN
     PREV199598097016
     Calcipotriol inhibits rectal epithelial cell proliferation in
TΙ
     ulcerative proctocolitis.
     Thopmas, M. G.; Nugent, K. P.; Forbes, A.; Williamson, R. C. N. (1)
ΑU
     (1) Royal Postgraduate Med. Sch., Hammersmith Hosp., Du Cane Rd., London,
CS
     W12 ONN UK
     Gut, (1994) Vol. 35, No. 12, pp. 1718-1720.
SO
     ISSN: 0017-5749.
DT
     Article
T.A
     English
    Vitamin D-3 reduces human rectal crypt cell production
AΒ
     rate (CCPR) and may thereby protect against colorectal cancer. Cell
     turnover is increased in ulcerative proctocolitis,
     which might therefore respond to vitamin D-3
    metabolites. This study investigated the effect of calcipotriol, a
     synthetic vitamin D-3 analogue that avoids
     hypercalcaemia, on human rectal CCPR in ulcerative
    proctocolitis. Paired rectal biopsy specimens from seven patients
     with severe disease were established in organ culture with or without
     calcipotriol (1 times 10-6 M). After 15 hours, vincristine (0.6 mu-g/ml)
     was added to induce metaphase arrest, and CCPR was determined by linear
     regression analysis of accumulated metaphases. Compared with values in 17
     controls with incidental anal conditions, median rectal CCPR was 28%
     higher in ulcerative proctocolitis: 5.90 (5.00-9.50) v
     4.80 (2.85-7.07) cells/crypt/hour (p lt 0.01). Calcipotriol reduced CCPR
     by 62% in patients with ulcerative proctocolitis, from
     5.90 (5.00-9.50) to 2.21 (0.81-3.22) cells/crypt/hour (median with range)
     p 1t 0.01. Thus calcipotriol can dampen the hyperproliferative state in
     ulcerative proctocolitis and could have a therapeutic
     role in the control of this inflammatory condition.
CC
     Cytology and Cytochemistry - Human *02508
     Biochemical Studies - General
                                      10063
     Biochemical Studies - Vitamins
     Biochemical Studies - Lipids
                                    10066
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Pathology, General and Miscellaneous - Therapy
                                                       12512
      Digestive System - Pathology *14006
     Endocrine System - General *17002
     Pharmacology - Clinical Pharmacology
     Pharmacology - Digestive System *22014
     Developmental Biology - Embryology - Morphogenesis, General *25508
BC
     Hominidae *86215
IT
     Major Concepts
        Cell Biology; Development; Endocrine System (Chemical Coordination and
        Homeostasis); Gastroenterology (Human Medicine, Medical Sciences);
        Pathology; Pharmacology
ΙT
     Chemicals & Biochemicals
        CALCIPOTRIOL; VITAMIN D3
     Miscellaneous Descriptors
TΤ
        CALCIPOTRIOL; GASTROINTESTINAL-DRUG; INFLAMMATION; VITAMIN D3
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        human (Hominidae)
ORGN Organism Superterms
        animals; chordates; humans; mammals; primates; vertebrates
     112965-21-6 (CALCIPOTRIOL)
RN
     67-97-0 (VITAMIN D3)
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L123 ANSWER 13 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

1994:106890 BIOSIS

DN PREV199497119890

- Longitudinal assessment of growth, mineral metabolism, and bone mass in ΤI pediatric Crohn's disease.
- Issenman, Robert M. (1); Atkinson, Stephanie A.; Radoja, Christine; ΑU Fraher, Laurence
- (1) Dep. Pediatrics, Children's Hosp. at Chedoke-McMaster, 1200 Main CS Street, Hamilton, ON L8N 3Z5 Canada
- Journal of Pediatric Gastroenterology and Nutrition, (1993) Vol. 17, No. SO 4, pp. 401-406. ISSN: 0277-2116.
- DTArticle
- English LA
- In children with inflammatory bowel disease, AB controversy continues about the use of long-term alternate day prednisone therapy (ADP) to suppress disease activity and to encourage appetite and growth. One possible side effect of both disease process and prednisone therapy is risk of development of osteoporosis. To evaluate this risk factor, growth, biochemical indices of mineral and vitamin D status, and bone mass were measured in nine adolescents with Crohn's disease (CD) who were treated with ADP (0.3 mg/kg gt 3 months per year) compared with eight adolescents treated with minimal ADP exposure ( 1t 3 months per year). Single photon densitometry was used to measure bone mineral mass at the 1/3 distal radius three times over 2 years. Mean age of the 17 CD boys was 13.9 +- 2.1 years at baseline. CD patients had lower bone BMC/BW mineral content/bone width (BMC/BW) compared with age- and height-matched normal boys at all times. The difference was less when compared to height-matched normal values as CD patients were shorter than healthy reference boys. Plasma 1,25-dihydroxyvitamin D, alkaline phosphatase, and parathyroid hormone significantly increased with treatment of disease but there were no differences between treatment groups. CD patients treated with ADP had similar heights and weights at baseline and demonstrated similar linear growth over 2 years (9.1 cm/2 years) to CD patients without ADP (10.3 cm/2 years). In both groups, BMC/BW increased significantly from year 1 to year 2, but absolute values for bone mass did not differ between the groups. These data suggest that over a 2-year treatment period male CD patients with chronic low-dose ADP exposure achieve linear growth rates and maintain bone mineralization at least as well as male CD patients who do
- Clinical Biochemistry; General Methods and Applications 10006 CC Biochemical Studies - Vitamins 10063

Biochemical Studies - Proteins, Peptides and Amino Acids 10064

Biochemical Studies - Sterols and Steroids

Biochemical Studies - Minerals 10069

Enzymes - Physiological Studies \*10808

Pathology, General and Miscellaneous - Inflammation and Inflammatory \*12508

Pathology, General and Miscellaneous - Therapy \*12512

Metabolism - Minerals \*13010

not receive ADP.

Metabolism - Fat-Soluble Vitamins \*13016

Nutrition - Fat-Soluble Vitamins \*13208

Digestive System - Pathology \*14006

Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies 15002

Endocrine System - Adrenals \*17004

Endocrine System - Parathyroid \*17010

Bones, Joints, Fasciae, Connective and Adipose Tissue - Physiology and Biochemistry \*18004

Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology \*18006

Pharmacology - Clinical Pharmacology \*22005

Pharmacology - Endocrine System \*22016

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Pharmacology - Immunological Processes and Allergy *22018
    Pediatrics *25000
    Developmental Biology - Embryology - Morphogenesis, General *25508
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
    Hominidae *86215
BC
    Major Concepts
ΙT
        Clinical Immunology (Human Medicine, Medical Sciences); Development;
        Endocrine System (Chemical Coordination and Homeostasis); Enzymology
        (Biochemistry and Molecular Biophysics); Gastroenterology (Human
        Medicine, Medical Sciences); Metabolism; Nutrition; Pathology;
        Pediatrics (Human Medicine, Medical Sciences); Pharmacology; Skeletal
        System (Movement and Support)
IT
    Chemicals & Biochemicals
        PREDNISONE; 1,25-DIHYDROXYVITAMIN D; VITAMIN D;
        ALKALINE PHOSPHATASE
    Miscellaneous Descriptors
ΙT
        ALKALINE PHOSPHATASE; ALTERNATE DAY PREDNISONE THERAPY; HORMONE-DRUG;
        HUMAN ADOLESCENT; IMMUNOSUPPRESSANT-DRUG; INFLAMMATORY
        BOWEL DISEASE; OSTEOPOROSIS; PARATHYROID HORMONE;
        PREDNISONE; VITAMIN D; 1,25-DIHYDROXYVITAMIN D
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        Hominidae (Hominidae)
ORGN Organism Superterms
        animals; chordates; humans; mammals; primates; vertebrates
RN
     53-03-2 (PREDNISONE)
     32222-06-3Q (1,25-DIHYDROXYVITAMIN D)
     66772-14-3Q (1,25-DIHYDROXYVITAMIN D)
       1406-16-2 (VITAMIN D)
     9001-78-9 (ALKALINE PHOSPHATASE)
L123 ANSWER 14 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
    1993:482091 BIOSIS
AN
     PREV199396115691
DN
    A study on interleukin-6 in inflammatory bowel
TΙ
     disease.
ΑU
     Yamakawa, Masaki
     Second Dep. Intern. Med., Nagasaki Univ. Sch. Med. Japan
CS
     Japanese Journal of Gastroenterology, (1993) Vol. 90, No. 6, pp.
SO
     1481-1488.
     ISSN: 0446-6586.
    Article
DT
LA
     Japanese
     Japanese; English
SL
    The production of interleukin-6 (IL-6) in patients with
     inflammatory bowel disease (IBD) has
     been measured, including the effects of steroid hormone,
     salicylazosulfapyridine (SASP) and its metabolites. In active
     Crohn's disease (CD) (n=12) and ulcerative
     colitis (UC) (n=9), rate of IL-6 positive group in serum was
     significantly higher than that in controls (n=20) (p lt 0.01, p lt 0.01).
     In active CD (n=9) and UC (n=9), the level of IL-6 production by
     peripheral blood mononuclear cells (PBMNC) was 22.8 +- 15.1ng/ml, 24.3 +-
     14.4ng/ml, and it was significantly higher than that in controls (n=15,
     8.0 + -6.6ng/ml (p lt 0.05, p lt 0.01). IL-6 production by PBMNC always
     showed the time dependent increase both in IBD and controls, and
     the level of IL-6 was always higher in IBD than that in controls
     during the culture time. Furthermore, IL-6 production by monocyte in UC
     (n=6, 4.4 +- 1.4 ng/ml) was significantly higher than that in controls
     (n=6, 1.7 +- 0.8 \text{ng/ml}) (p lt 0.01). The effects of steroid hormone, SASP
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and its metabolites on IL-6 production were also investigated. Steroid

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hormone significantly reduced IL-6 production by PBMNC, but others had no
     effect on IL-6 production. This study suggested that IL-6 might be
     involved in the pathophysiology of IBD.
    Cytology and Cytochemistry - Animal *02506
     Biochemical Studies - General
                                     10060
     Biochemical Studies - Proteins, Peptides and Amino Acids
                                                                10064
     Biochemical Studies - Sterols and Steroids
     Pathology, General and Miscellaneous - Therapy
                                                       12512
      Digestive System - Pathology *14006
     Endocrine System - General *17002
     Pharmacology - Digestive System *22014
ВC
     Hominidae *86215
    Major Concepts
IT
        Cell Biology; Endocrine System (Chemical Coordination and Homeostasis);
        Gastroenterology (Human Medicine, Medical Sciences); Pharmacology
     Chemicals & Biochemicals
IT
        SALICYLAZOSULFAPYRIDINE
ΙT
    Miscellaneous Descriptors
        CALBINDIN; CALCIUM ABSORPTION; HORMONE-DRUG; MESSENGER RNA; PARATHYROID
        HORMONE; VITAMIN D ANALOG; VITAMIN
       D RECEPTOR
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia; Muridae:
        Rodentia, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        rat (Muridae); Hominidae (Hominidae)
ORGN Organism Superterms
        animals; chordates; humans; mammals; nonhuman mammals; nonhuman
        vertebrates; primates; rodents; vertebrates
     599-79-1 (SALICYLAZOSULFAPYRIDINE)
RN
L123 ANSWER 15 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1993:333275 BIOSIS
ΑN
     PREV199345028000
DN
     Densitometric and biologic evaluation of bone status in patients with
ΤI
     ileo-anal pouch anastomosis (IAPA.
     Abitbol, V.; Chaussade, S.; Roux, C.; Pelleter, O.; Pigot, F.; Guillemant,
ΑU
     S.; Valleur, P.; Hautefeuille, P.; Amor, B.; et al.
     Service de Gastroenterol. Rhumatol., Hopital Cochin, Paris France
CS
     Gastroenterology, (1993) Vol. 104, No. 4 SUPPL., pp. A657.
SO
     Meeting Info.: 94th Annual Meeting of the American Gastroenterological
     Association Boston, Massachusetts, USA May 15-21, 1993
     ISSN: 0016-5085.
DT
     Conference
LA
     English
     General Biology - Symposia, Transactions and Proceedings of Conferences,
CC
     Congresses, Review Annuals
                                  00520
     Biochemical Studies - Vitamins
                                      10063
     Biochemical Studies - Minerals
                                      10069
     Anatomy and Histology, General and Comparative - Surgery *11105
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Metabolism - Minerals *13010
     Metabolism - Fat-Soluble Vitamins *13016
     Metabolism - Metabolic Disorders *13020
     Nutrition - Fat-Soluble Vitamins *13208
     Digestive System - Physiology and Biochemistry *14004
       Digestive System - Pathology *14006
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Physiology and
     Biochemistry *18004
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Hominidae *86215
BC
```

ΙT

Major Concepts

Digestive System (Ingestion and Assimilation); Gastroenterology (Human Medicine, Medical Sciences); Metabolism; Nutrition; Pathology; Skeletal System (Movement and Support); Surgery (Medical Sciences) Chemicals & Biochemicals IT VITAMIN D IT Miscellaneous Descriptors ABSTRACT; BONE MINERAL DENSITY; INFLAMMATORY BOWEL . DISEASE; VITAMIN D DEFICIENCY ORGN Super Taxa Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name Hominidae (Hominidae) ORGN Organism Superterms animals; chordates; humans; mammals; primates; vertebrates 1406-16-2 (VITAMIN D) L123 ANSWER 16 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 1993:241234 BIOSIS DN PREV199344114434 Prevalence of osteoporosis in patients with inflammatory ΤI bowel disease. Bjarnason, I. (1); MacPherson, A. J.; Buxton-Thomas, M.; Forgacs, I.; ΑU Moniz, C. (1) Dep. Clinical Biochem., King's Coll. Sch. Med., London SE5 9PJ UK CS Calcified Tissue International, (1993) Vol. 52, No. SUPPL. 1, pp. S65. SO Meeting Info.: XXIIIrd European Symposium on Calcified Tissues Heidelberg, Germany April 25-29, 1993 ISSN: 0171-967X. DTConference LA English General Biology - Symposia, Transactions and Proceedings of Conferences, CC Congresses, Review Annuals 00520 Mathematical Biology and Statistical Methods 04500 Biochemical Studies - Vitamins 10063 Biochemical Studies - Sterols and Steroids 10067 10069 Biochemical Studies - Minerals Chordate Body Regions - Back and Buttocks 11310 Pathology, General and Miscellaneous - Inflammation and Inflammatory Disease \*12508 Metabolism - Sterols and Steroids \*13008 Metabolism - Minerals \*13010 Metabolism - Fat-Soluble Vitamins \*13016 Metabolism - Metabolic Disorders \*13020 Nutrition - Malnutrition; Obesity \*13203 Nutrition - Minerals \*13206 Nutrition - Fat-Soluble Vitamins \*13208 Nutrition - General Dietary Studies \*13214 \*13226 Nutrition - Sterols and Steroids Digestive System - Pathology \*14006 Endocrine System - General \*17002 Endocrine System - Adrenals \*17004 Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology \*18006 Public Health - Public Health Administration and Statistics \*37010 Public Health: Epidemiology - Organic Diseases and Neoplasms \*37054 Hominidae \*86215 BC ΙT Major Concepts Endocrine System (Chemical Coordination and Homeostasis); Epidemiology (Population Studies); Gastroenterology (Human Medicine, Medical Sciences); Metabolism; Nutrition; Pathology; Public Health (Allied Medical Sciences); Skeletal System (Movement and Support) Chemicals & Biochemicals ΙT VITAMIN D; CALCIUM

Miscellaneous Descriptors

IT

ABSTRACT; CALCIUM; CORTICOSTEROIDS; CROHN'S DISEASE; ULCERATIVE COLITIS; VERTEBRAL BONE DENSITY; VITAMIN D

ORGN Super Taxa

Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia

ORGN Organism Name

human (Hominidae)

ORGN Organism Superterms

animals; chordates; humans; mammals; primates; vertebrates

RN 1406-16-2 (VITAMIN D) 7440-70-2 (CALCIUM)

L123 ANSWER 17 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

AN 1992:31034 BIOSIS

DN BA93:20309

TI NUTRITIONAL STATUS OF PATIENTS UNDERGOING ILEAL POUCH-ANAL ANASTOMOSIS.

AU PIRONI L; MIGLIOLI M; RUGGERI E; DALLASTA M A; POGGIOLI G; CAUDARELLA R; PIAZZI S; MINIERO R; GOZZETTI G; BARBARA L

CS IST. CLINICA MED. GASTROENTEROLOGIA, VIA MASSARENTI 9, 40138 BOLOGNA, ITALY.

SO CLIN NUTR (EDINB), (1991) 10 (5), 292-297. CODEN: CLNUDP. ISSN: 0261-5614.

FS BA; OLD

LA English

- The nutritional consequences of total colectomy and ileal pouch-anal AB anastomosis (IPAA) were assessed by evaluating 36 patients at the end of the defunctionalised stage (DS group) and 18 patients with recanalised IPAA (IPAA group). The changes in protein-calorie and zinc status occurring after the closure of the diverting ileostomy were evaluated also in 11 patients assessed both during the DS and the IPAA stage. The results were compared with those observed in 14 patients who underwent a Brooke-type permanent ileostomy (PI group). In the DS group there were protein-calorie malnutrition in 50% of cases characterised by body weight, TSF and AMC values lower than normal associated with normal serum protein levels; severe salt and water depletion with secondary aldosteronism in 90%; normal calcium-phosphorus balance in all but a few cases, low values of parameters related to vitamin D and K, Fe, Zn and Cu status in 6-25% of cases and normal folate status. In the IPAA group all the anthropmetric parameters improved significantly after the closure of the protective ileostomy, but muscle mass (AMC) remained lower than normal in 40% of cases; mild salt depletion (urinary Na/K ratio between 1 and 2) was observed in 1/3 of cases and of severe degree (urinary Na/K < 1) in 20%; lower serum Zn occurred in 60% of patients probably due to greater requirements of the metal, secondary to increased muscle protein synthesis; parameters of calcium-phosphorus balance, vitamin D and K, folate, Fe and Cu status, were normal in almost all the cases. In the PI group, protein-calorie and salt and mineral nutritional status were similar to those of the IPAA group, whereas Zn status was normal in all the patients and erythrocytes folate levels and prothrombin time were significantly lower than in the IPAA group. These last two results might be explained by the different characteristics of the small bowel flora occurring in the two types of ileostomy.
- CC Cytology and Cytochemistry Human 02508
  Mathematical Biology and Statistical Methods 04500
  Biochemistry Physiological Water Studies \*10011
  Biochemical Studies Vitamins 10063
  Biochemical Studies Proteins, Peptides and Amino Acids 10064
  Biochemical Studies Sterols and Steroids 10067
  Biochemical Studies Minerals 10069
  Anatomy and Histology, General and Comparative Surgery \*11105
  Pathology, General and Miscellaneous Inflammation and Inflammatory
  Disease \*12508

Pathology, General and Miscellaneous - Therapy 12512

```
Metabolism - Energy and Respiratory Metabolism *13003
    Metabolism - Sterols and Steroids *13008
    Metabolism - Minerals
                           *13010
    Metabolism - Fat-Soluble Vitamins *13016
    Metabolism - Water-Soluble Vitamins *13018
    Metabolism - Metabolic Disorders *13020
    Nutrition - Malnutrition; Obesity *13203
    Nutrition - Minerals *13206
    Nutrition - Fat-Soluble Vitamins *13208
    Nutrition - Water-Soluble Vitamins *13210
                                       *13226
    Nutrition - Sterols and Steroids
    Digestive System - General; Methods 14001
      Digestive System - Pathology *14006
    Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies
     *15002
    Blood, Blood-Forming Organs and Body Fluids - Blood Cell Studies *15004
    Endocrine System - Adrenals *17004
    Medical and Clinical Microbiology - General; Methods and Techniques 36001
    Microorganisms - Unspecified 01000
BC
    Hominidae 86215
    Miscellaneous Descriptors
ΙT
        HUMAN IRON ZINC COPPER POTASSIUM SODIUM VITAMIN D
        VITAMIN K FOLATE CALCIUM PHOSPHOROUS BALANCE WATER DEPLETION
        ERYTHROCYTES PROTHROMBIN TIME SECONDARY ALDOSTERONISM PROTEIN-CALORIE
        MALNUTRITION ULCERATIVE COLITIS FAMILIAL POLYPOSIS
        SMALL BOWEL FLORA METHOD BROOKE-TYPE PERMANENT ILEOSTOMY STATISTICS
    59-30-3 (FOLATE)
RN
       1406-16-2 (VITAMIN D)
     7439-89-6 (IRON)
     7440-09-7 (POTASSIUM)
    7440-23-5 (SODIUM)
     7440-50-8 (COPPER)
     7440-66-6 (ZINC)
     7440-70-2 (CALCIUM)
     9001-26-7 (PROTHROMBIN)
     12001-79-5 (VITAMIN K)
L123 ANSWER 18 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
    1989:416396 BIOSIS
AN
    BR37:71859
DN
    VITAMIN D METABOLISM BY MONOCYTES FROM PATIENTS WITH
ΤI
     INFLAMMATORY BOWEL DISEASE.
     SOSKOLNE W A; OFFENBACHER S; VAN DYKE T E
ΑŲ
     EMORY UNIV., ATLANTA, GA. USA.
ÇS
     67TH GENERAL SESSION OF THE INTERNATIONAL ASSOCIATION FOR DENTAL RESEARCH
SO
     (IADR), 6TH MEETING OF THE IADR IRISH DIVISION, 72ND ANNUAL MEETING OF THE
     SCANDINAVIAN ASSOCIATION FOR DENTAL RESEARCH AND THE 26TH ANNUAL MEETING
     OF THE CONTINENTAL EUROPEAN DIVISION OF THE IADR, DUBLIN, IRELAND, JUNE
     28-JULY 1, 1989. J DENT RES. (1989) 68 (SPEC ISSUE JUNE), 1006.
     CODEN: JDREAF. ISSN: 0022-0345.
DT
     Conference
FS
     BR; OLD
LA
     English
     General Biology - Symposia, Transactions and Proceedings of Conferences,
CC
     Congresses, Review Annuals 00520
     Cytology and Cytochemistry - Human 02508
     Biochemical Studies - Vitamins 10063
     Biochemical Studies - Sterols and Steroids 10067
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease 12508
     Metabolism - Fat-Soluble Vitamins *13016
       Digestive System - Pathology *14006
     Blood, Blood-Forming Organs and Body Fluids - Blood Cell Studies *15004
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Blood, Blood-Forming Organs and Body Fluids - Lymphatic Tissue and Reticuloendothelial System \*15008 Immunology and Immunochemistry - Immunopathology, Tissue Immunology \*34508 BC Hominidae 86215 Miscellaneous Descriptors IΤ ABSTRACT VITAMIN D 25 HYDROXYVITAMIN D-3 IMMUNOLOGY 1406-16-2 (VITAMIN D) RN 19356-17-3 (25 HYDROXYVITAMIN D-3) L123 ANSWER 19 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 1989:117076 BIOSIS AN BR36:62492 DN CHANGES OF THE CALCIUM METABOLISM IN INFLAMMATORY BOWEL TIDISEASES. KOCIAN J; KOCIANOVA J ΑU FAC. HOSP. BULOVKA, PRAGUE. CS GOEBELL, H., B. M. PESKAR AND H. MALCHOW (ED.). FALK SYMPOSIUM, 46. SO INFLAMMATORY BOWEL DISEASES: BASIC RESEARCH AND CLINICAL IMPLICATIONS; TITISEE, WEST GERMANY, JUNE 7-9, 1987. XVIII+449P. KLUWER ACADEMIC PUBLISHERS: DORDRECHT, NETHERLANDS; BOSTON, MASSACHUSETTS, USA. ILLUS. (1988) 0 (0), 417. CODEN: FASYDI. ISSN: 0161-5580. ISBN: 0-7462-0067-6. DT Conference FS BR; OLD LAEnglish General Biology - Symposia, Transactions and Proceedings of Conferences, CC Congresses, Review Annuals 00520 Biochemical Studies - Vitamins 10063 Biochemical Studies - Sterols and Steroids 10067 Biochemical Studies - Carbohydrates 10068 Biochemical Studies - Minerals 10069 Pathology, General and Miscellaneous - Inflammation and Inflammatory Disease 12508 Metabolism - Carbohydrates \*13004 Metabolism - Minerals \*13010 Metabolism - Fat-Soluble Vitamins \*13016 Metabolism - Metabolic Disorders \*13020 Digestive System - Pathology \*14006 Endocrine System - Adrenals 17004 Bones, Joints, Fasciae, Connective and Adipose Tissue - Physiology and Biochemistry \*18004 Pharmacology - Clinical Pharmacology \*22005 Pharmacology - Digestive System \*22014 Pharmacology - Endocrine System \*22016 Immunology and Immunochemistry - Immunopathology, Tissue Immunology \*34508 BC Hominidae 86215 IT Miscellaneous Descriptors ABSTRACT HUMAN CORTICOSTEROID TREATMENT VITAMIN D METABOLISM BONE MINERALIZATION CROHN'S DISEASE ULCERATIVE COLITIS INTESTINAL WALL AFFECTION LACTOSE INTOLERANCE 63-42-3 (LACTOSE) 1406-16-2 (VITAMIN D) 7440-70-2 (CALCIUM) L123 ANSWER 20 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. ΑN 1988:367178 BIOSIS DN BR35:51791 DECREASED BONE MINERALIZATION IN PATIENTS WITH INFLAMMATORY TΙ BOWEL DISEASE IBD. STALLMACH A; FELSENBERG D; PIONTEK A; VALLO M; ZEITZ M; RIECKEN E O

ΑU

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DEP. MED., FREE UNIV. BERLIN, WEST BERLIN.
CS
    89TH ANNUAL MEETING OF THE AMERICAN GASTROENTEROLOGICAL ASSOCIATION, NEW
SO
    ORLEANS, LOUISIANA, USA, MAY 14-20, 1988. GASTROENTEROLOGY. (1988) 94 (5
     PART 2), A440.
    CODEN: GASTAB. ISSN: 0016-5085.
    Conference
DT
FS
    BR; OLD
    English
LA
    General Biology - Symposia, Transactions and Proceedings of Conferences,
CC
    Congresses, Review Annuals 00520
     Biochemical Studies - Vitamins 10063
     Biochemical Studies - Minerals 10069
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease 12508
    Metabolism - Minerals *13010
    Metabolism - Fat-Soluble Vitamins *13016
     Nutrition - Malnutrition; Obesity *13203
       Digestive System - Pathology *14006
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology 34508
BC
     Hominidae 86215
ΙT
    Miscellaneous Descriptors
        ABSTRACT HUMAN CROHN'S DISEASE ULCERATIVE
        COLITIS CALCIUM VITAMIN D
     1406-16-2 (VITAMIN D)
RN
     7440-70-2 (CALCIUM)
L123 ANSWER 21 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
    1988:219639 BIOSIS
ΑN
DN
     BA85:108874
     CARIES RISK IN PATIENTS WITH CROHN'S DISEASE A PILOT STUDY.
ΤI
     BEVENIUS J
ΑU
     DEP. CARIOL., ODONTOL. FAK., ODONTOL. KLINIKERNA, BOX 4064, S-141 04
CS
     HUDDINGE, SWEDEN.
     ORAL SURG ORAL MED ORAL PATHOL, (1988) 65 (3), 304-307.
SO
     CODEN: OSOMAE. ISSN: 0030-4220.
FS
    BA; OLD
    English
T.A
    Crohn's disease is a chronic inflammatory
AB
    bowel disease of unknown cause with unpredictable
     remissions and exacerbations. Associated nutritional deficiences include
     those involving zinc, magnesium, vitamin B12, folic acid, and
     vitamin D. A group of patients with Crohn's
     disease underwent detailed cariologic investigation at the Department of
     Cariology, Karolinska Institutet, Stockholm [Sweden]. Factors predisposing
     to caries were evaluated according to Krasse's concept of caries risk. On
     this basis, 11 of the 15 patients had a high caries risk. The concept of
     caries risk acknowledges the multifactorial background of caries
     initiation and progression and, in this pilot study, has proved to be an
     appropriate basis for evaluation of patients with chronic disease.
     Guidelines for preventive programs appropriate for patients with
     Crohn's disease, based on the findings of this study, are
     presented.
     Biochemical Studies - Vitamins 10063
CC
     Biochemical Studies - Sterols and Steroids 10067
     Biochemical Studies - Minerals 10069
     Pathology, General and Miscellaneous - Diagnostic *12504
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Nutrition - Malnutrition; Obesity *13203
     Nutrition - Minerals *13206
     Nutrition - Fat-Soluble Vitamins *13208
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Nutrition - Water-Soluble Vitamins \*13210

Digestive System - Pathology \*14006 Dental and Oral Biology - Pathology \*19006 Medical and Clinical Microbiology - Bacteriology \*36002 Public Health - Public Health Administration and Statistics \*37010 BC Hominidae 86215 Miscellaneous Descriptors IT SWEDEN NUTRITIONAL DEFICIENCIES ZINC MAGNESIUM VITAMIN B-12 FOLIC ACID VITAMIN D CHRONIC INFLAMMATORY BOWEL DISEASE PREDISPOSING FACTORS 59-30-3 (FOLIC ACID) RN 68-19-9 (VITAMIN B-12) 1406-16-2 (VITAMIN D) 7439-95-4 (MAGNESIUM) 7440-66-6 (ZINC) L123 ANSWER 22 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 1987:314049 BIOSIS ΑN BA84:33556 DN THE ROLE OF NUTRITION IN THE TREATMENT OF INFLAMMATORY TΤ BOWEL DISEASE. ΑU MERYN S I. UNIVERSITAETSKLIN. GASTROENTEROL. HEPATOL., LAZARETTGASSE 14, A-1090 CS WIEN, AUSTRIA. WIEN KLIN WOCHENSCHR, (1986 (RECD 1987)) 98 (22), 774-779. SO CODEN: WKWOAO. ISSN: 0043-5325. BA; OLD FS LAGerman The clinical picture and course of inflammatory bowel AB disease are influenced by nutritional abnormalities and malnutrition. Interest at present concentrates on high-fibre low-refined sugar diets, elimination diets with identification of specific food intolerance and low-residue diets. All three failed to show significant positive effects on the course of the disease, need for hospitalisation, surgical procedures required or postoperative recurrence. Only a low lactose diet seems to be justified, since we found lactose intolerance in 25-35% of patients with inflammatory bowel disease, as compared with 5-10% in the normal population. In 25 patients with Crohn's disease (CD) a reduction in inflammatory activity and improvement of nutritional status was obtained with parenteral nutrition (PN). Nevertheless, longer follow up periods revealed no additional benefit in comparison with conventional therapies. Furthermore, the combination of PN and total bowel rest resulted in the same improvement as with PN alone. 25 patients with CD manifesting an. acute phase of the condition were treated with tube feeding (TF) as primary therapy. TF reduced CD activity and improved nutritional status in 15 patients with small bowel disease, whereas the patients with colonic disease and extraintestinal manifestations did not react. A comparison of the effect of PN and TF in 10 patients with CD showed no significant difference with regard to clinical course and objective parameters. In view of the high costs and risks of complications of PN, TF is recommended as primary therapy for the acute phase of CD. The importance of substitution therapy, especially of vitamin D, is documented. Biochemical Studies - Carbohydrates 10068 CC Pathology, General and Miscellaneous - Diagnostic 12504 Pathology, General and Miscellaneous - Inflammation and Inflammatory \*12508 Pathology, General and Miscellaneous - Therapy \*12512 Metabolism - Carbohydrates 13004 Metabolism - Metabolic Disorders 13020 Nutrition - General Studies, Nutritional Status and Methods \*13202 Nutrition - Malnutrition; Obesity \*13203

Nutrition - Prophylactic and Therapeutic Diets \*13218

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*13220
     Nutrition - Carbohydrates
     Digestive System - General; Methods *14001
       Digestive System - Pathology *14006
     Routes of Immunization, Infection and Therapy 22100
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
BC
     Hominidae 86215
    Miscellaneous Descriptors
IT
        HUMAN TUBE FEEDING CROHN'S DISEASE ULCERATIVE
        COLITIS PARENTERAL PATHOGENESIS FIBER SUGAR LACTOSE INTOLERANCE
     63-42-3Q, 16984-38-6Q (LACTOSE)
RN
L123 ANSWER 23 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1986:142962 BIOSIS
DN
     BA81:53378
    VITAMIN D STATUS IN CROHN'S DISEASE
TI
     ASSOCIATION WITH NUTRITION AND DISEASE ACTIVITY.
     HARRIES A D; BROWN R; HEATLEY R V; WILLIAMS L A; WOODHEAD S; RHODES J
ΑU
     UNIV. HOSP. WALES, HEATH PARK, CARDIFF.
CS
     GUT, (1985) 26 (11), 1197-1203.
SO
     CODEN: GUTTAK. ISSN: 0017-5749.
     BA; OLD
FS
LA
     English
     Forty patients with Crohn's disease were divided into
AΒ
     undernourished (18) and well nourished (22) groups depending on whether
     their midarm circumference was below or above 90% of the ideal standard.
     Plasma 25-(OH)D3 and the dihydroxylated metabolites, 24,25-(OH)2D3 and
     1,25-(OH)2D3 were measured in the summer. Results were related to clinical
     and biochemical parameters and also compared with results from patients
     with ulcerative colitis and healthy subjects who
     served as controls. Plasma 25-(OH)D3 was reduced in the undernourished
     Crohn's groups compared with the well nourished Crohn's
     group, who did not differ from the controls. Over 50% of the
     undernourished Crohn's group had evidence of secondary
     hyperparathyroidism and raised alkaline phosphatase concentrations,
     although concentrations of 1,25-(OH)2D3 were normal. The low 25-(OH)D3
     concentrations related to disease activity. It is suggested that
     undernourished Crohn's patients who have high levels of disease
     activity are at risk of vitamin D deficiency, and
     attempts should be made to improve their vitamin D
     nutrition.
     Clinical Biochemistry; General Methods and Applications 10006
     Biochemical Studies - Vitamins 10063
     Biophysics - General Biophysical Techniques 10504
     Enzymes - Physiological Studies *10808
     Pathology, General and Miscellaneous - Diagnostic 12504
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Metabolism - Fat-Soluble Vitamins *13016
     Metabolism - Metabolic Disorders *13020
     Nutrition - Malnutrition; Obesity *13203
Nutrition - Fat-Soluble Vitamins *13208
     Digestive System - General; Methods 14001
       Digestive System - Pathology *14006
     Endocrine System - Thyroid *17018
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
     Hominidae 86215
BC
     Miscellaneous Descriptors
        HUMAN 1 25 DIHYDROXYVITAMIN D-3 24 25 DIHYDROXYVITAMIN D-3 25
        HYDROXYVITAMIN D-3 MALNUTRITION DIHYDROXYLATED METABOLITE
        ULCERATIVE COLITIS ALKALINE PHOSPHATASE
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HYPERPARATHYROIDISM

```
1406-16-2 (VITAMIN D)
RN
     9001-78-9 (ALKALINE PHOSPHATASE)
     19356-17-3 (25 HYDROXYVITAMIN D-3)
     32222-06-3 (1 25 DIHYDROXYVITAMIN D-3)
     40013-87-4 (24 25 DIHYDROXYVITAMIN D-3)
L123 ANSWER 24 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1986:138335 BIOSIS
AN
DN
     BA81:48751
     BONE METABOLIC DISORDER DURING STEROID THERAPY FOR INFLAMMATORY
ΤI
     BOWEL DISEASES.
     TADA M; SHIMIZU S; KAWAI K; WATANABE Y
ΑU
     DEP. INTERNAL MED., KYOTO FIRST RED CROSS HOSPITAL, KYOTO, JAPAN.
CS
     J JPN SOC COLO-PROCTOL, (1985) 38 (6), 663-668.
SO
     CODEN: NDKGAU. ISSN: 0047-1801.
FS
     BA; OLD
     Japanese
LA
     Bone metabolic disorder is one of the untoward effects caused by steroid
ΑB
    administration for inflammatory bowel diseases
     . During steroid therapy, we tried to assess its effects on bone
     metabolism by means of the microdensitometry method. Using MCI,
     .DELTA.GSmin and .SIGMA.GS/D as indicators, the amount of administered
     prednisolone correlated with the degree of osteoporotic changes. Serum
     calcium, phosphorus, alkaline phosphatase and the N-terminal of PTH
     (parathormone) were also measured during the course, showing that the
     serum levels of calcium and phosphorus deviated in some cases where the
     doses of steroid were low. Administration of activated vitamin
     D (1.alpha.-OH-D3), 0.5
     .mu.g per day, during steroid therapy showed a tendency to prevent the
     development of osteoporosis and/or normalize the values already mentioned,
     in so far as the cumulative steroid dose was less than 4000 mg. These data
     indicated that, during steroid therapy, attention should be directed to
     its harmful effects on bone metabolism, and that the desirable effects of
     1.alpha.-OH-D3 should be
     appreciated.
     Cytology and Cytochemistry - Human 02508
     Biochemical Studies - Proteins, Peptides and Amino Acids 10064
     Biochemical Studies - Sterols and Steroids 10067
     Biochemical Studies - Minerals 10069
     Enzymes - General and Comparative Studies; Coenzymes *10802
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
              *12508
     Disease
     Pathology, General and Miscellaneous - Therapy
     Digestive System - General; Methods *14001
     Digestive System - Physiology and Biochemistry *14004
       Digestive System - Pathology *14006
     Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies
     Endocrine System - Adrenals *17004
     Endocrine System - Parathyroid *17010
     Bones, Joints, Fasciae, Connective and Adipose Tissue - General; Methods
     *18001
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Physiology and
     Biochemistry *18004
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Pharmacology - Drug Metabolism; Metabolic Stimulators *22003
     Pharmacology - Clinical Pharmacology
                                             *22005
     Pharmacology - Connective Tissue, Bone and Collagen - Acting Drugs 22012
     Pharmacology - Digestive System *22014
     Pharmacology - Endocrine System *22016
     Pharmacology - Immunological Processes and Allergy 22018
Toxicology - Pharmacological Toxicology *22504
     Toxicology - Antidotes and Preventative Toxicology *22505
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Immunology and Immunochemistry - Immunopathology, Tissue Immunology 34508
ВC
     Hominidae 86215
    Miscellaneous Descriptors
IT
        HUMAN PREDNISOLONE VITAMIN D HORMONE-DRUG
        VITAMIN-DRUG ANTIDOTE-DRUG PHARMACOTOXICITY MICRODENSITOMETRY
       OSTEOPOROSIS CALCIUM PHOSPHORUS ALKALINE PHOSPHATASE PARATHORMONE
     50-24-8 (PREDNISOLONE)
RN
       1406-16-2 (VITAMIN D)
     7440-70-2 (CALCIUM)
     7723-14-0 (PHOSPHORUS)
     9001-78-9 (ALKALINE PHOSPHATASE)
     9002-64-6 (PARATHORMONE)
L123 ANSWER 25 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1986:101564 BIOSIS
ΑN
     BA81:11980
DN
     VITAMIN D ABSORPTION IN HEALTHY SUBJECTS AND IN
TΤ
     PATIENTS WITH INTESTINAL MALABSORPTION SYNDROMES.
     LO C W; PARIS P W; CLEMENS T L; NOLAN J; HOLICK M F
ΑU
     USDA HUMAN NUTRITION RES. CENT., TUFTS UNIV., 711 WASHINGTON ST., BOSTON,
CS
     MASS. 02111.
     AM J CLIN NUTR, (1985) 42 (4), 644-649.
SO
     CODEN: AJCNAC. ISSN: 0002-9165.
FS
     BA; OLD
LA
     English
     We developed a test procedure for the clinical evaluation of the
AB
     absorption of vitamin D. Serum vitamin
     D concentrations were evaluated in seven patients with intestinal
     fat malabsorption syndromes and in seven healthy, normal subjects, after
     being given a single oral dose of 50,000 IU (1.25 mg) vitamin D2. In the
     normal subjects, serum vitamin D concentrations rose
     from a baseline of less than 5 ng/ml to a peak of over 50 ng/ml by 12 h,
     gradually falling to baseline levels by 3 days. In five of the seven
     patients with intestinal fat malabsorption, oral administration of 50,000
     IU vitamin D2 did not raise serum vitamin D
     concentrations above 10 ng/ml. Two patients with severe
     inflammatory bowel disease had a normal
     absorption pattern, however. These findings suggest that an oral
     vitamin D absorption test may be of value for
     determination of patients at risk for development of vitamin
     D deficiency. They also raise questions about the efficacy of oral
     vitamin D preparations in patients with intestinal fat
     malabsorption.
     Biochemical Studies - Vitamins 10063
CC
     Biochemical Studies - Sterols and Steroids 10067
     Pathology, General and Miscellaneous - Diagnostic 12504
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Metabolism - Fat-Soluble Vitamins *13016
     Nutrition - Malnutrition; Obesity *13203
     Nutrition - Fat-Soluble Vitamins *13208
     Digestive System - General; Methods 14001
     Digestive System - Physiology and Biochemistry *14004
       Digestive System - Pathology *14006
     Dental and Oral Biology - General; Methods 19001
     Routes of Immunization, Infection and Therapy 22100
BC
     Hominidae 86215
     Miscellaneous Descriptors
TT
          VITAMIN D DEFICIENCY INFLAMMATORY
        BOWEL DISEASE INTESTINAL FAT MALABSORPTION
RN
     1406-16-2 (VITAMIN D)
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L123 ANSWER 26 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

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qazi - 10 / 036819
     1985:242376 BIOSIS
ΑN
     BA79:22372
DN
     OSTEOPENIA WITH NORMAL VITAMIN D METABOLITES AFTER
ΤI
     SMALL-BOWEL RESECTION FOR CROHNS DISEASE.
     HESSOV I; MOSEKILDE L; MELSEN F; FASTH S; HULTEN L; LUND B; LUND B;
ΑU
     SORENSEN O H
     DEP. SURGERY I, AARHUS AMTSSYGEHUS, DK-8000 AARHUS C, DEN.
CS
     SCAND J GASTROENTEROL, (1984) 19 (5), 691-696.
SO
     CODEN: SJGRA4. ISSN: 0036-5521.
FS
     BA; OLD
     English
LA
     Unselected patients (36) were investigated 3-24 yr (mean, 7.8 yr) after
AB
     small-bowel resection for Crohn's disease (mean small intestinal
     resection, 105 cm). Iliac crest bone biopsies after in vivo tetracycline
     double-labeling showed a markedly reduced trabecular bone mass (controls,
     0.25 .+-. 0.06; patients, 0.15 .+-. 0.05; P < 0.01). The average bone
     remodeling and osteoid mineralization was normal, and only 2 patients
     demonstrated signs of frank but slight osteomalacia. The mean serum levels
     of the 3 vitamin D metabolites 25-hydroxyvitamin D,
     24,25-dihydroxyvitamin D and 1,25-dihydroxyvitamin D were normal. The
     observed reduction in trabecular bone mass may theoretically be followed
     by an increased risk of spontaneous fractures.
     Mathematical Biology and Statistical Methods 04500
     Biochemical Studies - Vitamins 10063
     Biochemical Studies - Minerals 10069
     Anatomy and Histology, General and Comparative - Surgery 11105
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease 12508
     Pathology, General and Miscellaneous - Therapy
                                                      12512
     Metabolism - Minerals *13010
     Metabolism - Fat-Soluble Vitamins *13016
       Digestive System - Pathology *14006
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
     Hominidae 86215
BC
     Miscellaneous Descriptors
IT
        HUMAN FRACTURE BONE REMODELING OSTEOID MINERALIZATION RISK
     1406-16-2 (VITAMIN D)
RN
L123 ANSWER 27 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1984:281296 BIOSIS
AN
     BA78:17776
DN
     A SURVEY OF VITAMIN D DEFICIENCY IN GASTRO INTESTINAL
ΤI
     AND LIVER DISORDERS.
     DIBBLE J B; SHERIDAN P; LOSOWSKY M S
ΑU
     DEP. MED., ST. JAMES UNIV. HOSP., LEEDS.
CS
     O J MED, (1984) 53 (209), 119-134.
SO
     CODEN: QJMEA7. ISSN: 0033-5622.
FS
     BA; OLD
     English
LΑ
     A survey of vitamin D status in 152 patients with
AB
     chronic gastrointestinal conditions and 104 patients with chronic liver
     diseases is presented. Mild deficiency was common and severe deficiency,
     as judged by plasma 25-OHD [25-hydroxy-vitamin D]
     levels < 8 nmol/1, was encountered in every disease category tested. In
     the gastrointestinal disease patients, deficiency was significantly more
     common in patients following gastroenterostomy than other gastric surgery,
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in patients with active Crohn's disease than in those with

features. Deficiency was as common in patients with Crohn's

inactive disease, and in patients with chronic pancreatitis or pancreatic carcinoma with cholestatic features than in those without cholestatic

disease who had not been treated surgically as in those who had. There was

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no significant correlation between plasma 25-OHD levels and any laboratory
    index of malabsorption or malnutrition except from serum albumin in the
    qastric surgery patients, Hb and ESR [erythrocyte sedimentation rate] in
    the Crohn's disease patients, and albumin and vitamin E in the
    group of patients with gastrointestinal disorders taken as a whole. In the
    chronic liver disease patients, those with late primary biliary cirrhosis
    had lower plasma 25-OHD levels than those with histological Stage I and II
    disease who all had normal levels, and those with pruritus and jaundice
    were more commonly severely deficient. Whatever the underlying disease
    process, patients with other coincidental medical conditions were much
    more likely to be deficient as were patients with cholestasis. Evidence of
    secondary hyperparathyroidism and osteomalacia on bone histology indicated
    the clinical relevance of the vitamin D deficiency.
    This study showed no relationship between abnormal plasma vitamin
    D binding protein levels and vitamin deficiency.
    Microscopy Techniques - Histology and Histochemistry 01056
    Cytology and Cytochemistry - Human 02508
    Clinical Biochemistry; General Methods and Applications
    Biochemical Studies - Vitamins 10063
    Biochemical Studies - Proteins, Peptides and Amino Acids 10064
    Biochemical Studies - Lipids 10066
    Biochemical Studies - Sterols and Steroids 10067
    Anatomy and Histology, General and Comparative - Surgery *11105
     Pathology, General and Miscellaneous - Comparative
    Pathology, General and Miscellaneous - Inflammation and Inflammatory
    Disease 12508
    Pathology, General and Miscellaneous - Therapy
    Metabolism - Lipids 13006
    Metabolism - Sterols and Steroids *13008
    Metabolism - Proteins, Peptides and Amino Acids *13012
    Metabolism - Porphyrins and Bile Pigments
    Metabolism - Fat-Soluble Vitamins *13016
    Metabolism - Metabolic Disorders *13020
    Nutrition - Malnutrition; Obesity *13203
                                      *13208
    Nutrition - Fat-Soluble Vitamins
    Nutrition - Pathogenic Diets *13216
     Digestive System - General; Methods 14001
      Digestive System - Pathology *14006
     Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies
    15002
    Blood, Blood-Forming Organs and Body Fluids - Blood Cell Studies *15004
    Endocrine System - Parathyroid *17010
    Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
    Hominidae 86215
    Miscellaneous Descriptors
       HUMAN ERYTHROCYTE SEDIMENTATION RATE ALBUMIN HEMO GLOBIN VITAMIN E 25
       HYDROXY VITAMIN D PRURITUS JAUNDICE CHOLESTASIS
       PRIMARY BILIARY CIRRHOSIS OSTEO MALACIA CROHNS
       DISEASE MAL ABSORPTION MAL NUTRITION HYPER PARATHYROIDISM
        GASTRO ENTEROSTOMY
    1406-16-2 (VITAMIN D)
    1406-18-4 (VITAMIN E)
     19356-17-3Q, 21343-40-8Q, 64719-49-9Q (25 HYDROXY VITAMIN
L123 ANSWER 28 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
    1983:290332 BIOSIS
    BA76:47824
    CALCIUM METABOLISM IN SUBJECTS LIVING WITH A PERMANENT ILEOSTOMY.
    KENNEDY H J; COMPSTON J; HEYNEN G; KANIS J A; MERRETT A L; TRUELOVE S C;
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BC

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RN

AN

DN

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ΑU

WARNER G T

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GASTROENTEROLOGY UNIT, RADCLIFFE INFIRMARY, OXFORD OX2 6HE, GB.
    DIGESTION, (1983) 26 (3), 131-136.
SO
    CODEN: DIGEBW. ISSN: 0012-2823.
FS
    BA; OLD
LA
    English
    Several indices of Ca metabolism were studied in 39 subjects living with a
AB
     permanent ileostomy after proctocolectomy for ulcerative
     colitis, and in a control group of 39 healthy volunteers, matched
     for age and sex. No significant differences were found in plasma levels of
     Ca, phosphate, Mg, parathyroid hormone, calcitonin and 25-hydroxy-
     vitamin D nor in the urinary excretion of Ca and
    phosphate, but the alkaline phosphatase was raised in the ileostomists.
     The bone density of ileostomists was rather low, but the difference from
     the control subjects was not statistically significant. The absorption of
     Ca was measured by means of a total body counter. The ileostomists
     retained significantly more Ca than expected. This may represent the
     correction of a state of Ca deficiency at the time of proctocolectomy, due
     to the effects of the colitis and its medical treatment with
     corticosteroids.
     Biochemical Studies - General 10060
     Biochemical Studies - Vitamins 10063
     Biochemical Studies - Proteins, Peptides and Amino Acids 10064
     Biochemical Studies - Sterols and Steroids 10067
     Biochemical Studies - Minerals 10069
     Biophysics - Membrane Phenomena 10508
     Enzymes - Physiological Studies *10808
     Anatomy and Histology, General and Comparative - Surgery *11105
     Physiology, General and Miscellaneous - Instrumentation 12004
               12100
     Movement
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
             12508
     Pathology, General and Miscellaneous - Therapy
     Metabolism - General Metabolism; Metabolic Pathways 13002
     Metabolism - Lipids *13006
     Metabolism - Minerals *13010
     Metabolism - Proteins, Peptides and Amino Acids *13012
     Nutrition - Malnutrition; Obesity *13203
     Digestive System - General; Methods *14001
       Digestive System - Pathology *14006
     Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies
     15002
     Urinary System and External Secretions - Physiology and Biochemistry
     15504
     Endocrine System - Adrenals *17004
     Endocrine System - Parathyroid *17010
     Endocrine System - Thyroid *17018
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Physiology and
     Biochemistry 18004
     Pharmacology - Clinical Pharmacology
                                            22005
     Pharmacology - Digestive System *22014
     Hominidae 86215
BC
     Miscellaneous Descriptors
TΤ
        CORTICO STEROIDS HORMONE-DRUG ANTIINFLAMMATORY
        GASTROINTESTINAL-DRUG ULCERATIVE COLITIS CALCIUM
        DEFICIENCY PARATHYROID HORMONE CALCITONIN 25 HYDROXY VITAMIN
        D PROCTO COLECTOMY ALKALINE PHOSPHATASE PHOSPHATE MAGNESIUM
        URINARY EXCRETION BONE DENSITY ABSORPTION
     7439-95-4 (MAGNESIUM)
RN
     7440-70-2 (CALCIUM)
     9001-78-9 (ALKALINE PHOSPHATASE)
     9007-12-9 (CALCITONIN)
     14265-44-2 (PHOSPHATE)
     19356-17-3Q, 21343-40-8Q, 64719-49-9Q (25 HYDROXY VITAMIN
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D)

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L123 ANSWER 29 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1983:228742 BIOSIS
     BA75:78742
DN
     VITAMIN D DEFICIENCY AND BONE DISEASE IN PATIENTS WITH
TΙ
     CROHNS DISEASE.
     DRISCOLL R H JR; MEREDITH S C; SITRIN M; ROSENBERG I H
ΑU
     UNIV. CHICAGO, 950 EAST 59TH ST., BOX 400, CHICAGO, ILL. 60637.
CS
     GASTROENTEROLOGY, (1982) 83 (6), 1252-1258.
SO
     CODEN: GASTAB. ISSN: 0016-5085.
     BA; OLD
FS
LA
     English
     The prevalence of vitamin D deficiency in
AΒ
     Crohn's disease and the relationship of vitamin
     D status to metabolic bone disease have not been fully
     characterized. Serum 25-hydroxyvitamin D was measured in 82 patients with
     Crohn's disease: 65% of Crohn's disease patients had a
     low serum 25-hydroxyvitamin D concentration; 25% had deficient levels (<
     10 ng/ml). The lowest 25-hydroxyvitamin D levels were observed in patients
     with previous ileal resections. Nine patients were studied in detail
     including transiliac needle bone biopsies; 6 had osteomalacia and 3
     osteoporosis. Six patients had repeat bone biopsies 9-18 mo. after
     vitamin D treatment. Three patients with osteomalacia
     and low serum 25-hydroxyvitamin D levels showed histologic improvement
     after therapy with oral vitamin D restored serum
     25-hydroxyvitamin D levels to normal. The adequacy of therapy was assessed
     accurately by monitoring serum 25-hydroxyvitamin D concentration. Three
     patients with metabolic bone disease with normal serum 25-hydroxyvitamin D
     levels at diagnosis did not show histologic improvement after receiving
     vitamin D.
     Microscopy Techniques - Histology and Histochemistry 01056
CC
     Biochemical Studies - Vitamins 10063
     Biochemical Studies - Sterols and Steroids 10067
     Biochemical Studies - Minerals 10069
     Anatomy and Histology, General and Comparative - Surgery 11105
     Anatomy and Histology, General and Comparative - Microscopic and
     Ultramicroscopic Anatomy 11108
     Pathology, General and Miscellaneous - Diagnostic 12504
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Pathology, General and Miscellaneous - Therapy
                                                       12512
     Metabolism - Minerals *13010
     Metabolism - Fat-Soluble Vitamins *13016
     Nutrition - Malnutrition; Obesity *13203
Nutrition - Fat-Soluble Vitamins *13208
     Nutrition - Prophylactic and Therapeutic Diets *13218
     Digestive System - General; Methods 14001
       Digestive System - Pathology *14006
     Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies
     15002
     Bones, Joints, Fasciae, Connective and Adipose Tissue - General; Methods
     18001
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Anatomy 18002
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Dental and Oral Biology - General; Methods 19001
     Routes of Immunization, Infection and Therapy 22100
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
BC
     Hominidae 86215
     Miscellaneous Descriptors
ΙT
        METABOLIC BONE DISEASE OSTEO MALACIA OSTEO POROSIS ILEAL RESECTION
     1406-16-2 (VITAMIN D)
RN
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L123 ANSWER 30 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1983:120003 BIOSIS
ΑN
     BR25:45003
DN
     BONE DISEASE AND HEPATO BILIARY DISORDERS.
ΤI
     JUTTMAN J R
ΑU
     DEP. MED. III, HOSP. DIJKZIGT, ERASMUS UNIV., ROTTERDAM.
CS
     LEENDERT SCHALM SYMPOSIUM ON PRIMARY BILIARY CIRRHOSIS HELD AT THE MEETING
SO
     OF THE NETHERLANDS ASSOCIATION FOR THE STUDY OF THE LIVER, MAY 11, 1982.
     NETH J MED. (1982) 25 (8), 290.
     CODEN: NLJMAV. ISSN: 0300-2977.
DT
     Conference
     BR; OLD
FS
     English
LA
     General Biology - Symposia, Transactions and Proceedings of Conferences,
CC
     Congresses, Review Annuals 00520
     Biochemical Studies - Vitamins 10063
     Biochemical Studies - Proteins, Peptides and Amino Acids 10064
     Biochemical Studies - Lipids 10066
     Biochemical Studies - Sterols and Steroids 10067
     Biochemical Studies - Minerals 10069
     Anatomy and Histology, General and Comparative - Surgery 11105
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease 12508
     Metabolism - Lipids 13006
     Metabolism - Minerals *13010
     Metabolism - Proteins, Peptides and Amino Acids 13012
     Metabolism - Fat-Soluble Vitamins *13016
     Nutrition - Malnutrition; Obesity *13203
     Nutrition - Pathogenic Diets 13216
     Nutrition - Proteins, Peptides and Amino Acids
                                                      13224
     Digestive System - General; Methods 14001
       Digestive System - Pathology *14006
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
     Plant Physiology, Biochemistry and Biophysics - Chemical Constituents
     51522
     Gramineae 25305
BC
     Hominidae 86215
     Miscellaneous Descriptors
IT
        ABSTRACT HUMAN VITAMIN D METABOLISM DISTURBANCE
        CALCIUM METABOLISM DISTURBANCE OSTEO MALACIA CROHNS
        DISEASE CELIAC DISEASE PANCREATIC INSUFFICIENCY PRIMARY BILIARY
        CIRRHOSIS CHOLESTATIC LIVER DISEASE OSTEO POROSIS GASTRECTOMY JEJUNO
        ILEAL BYPASS
RN
     1406-16-2 (VITAMIN D)
     7440-70-2 (CALCIUM)
L123 ANSWER 31 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1981:13534 BIOSIS
ΑN
     BR20:13534
DN
     CALCIUM ABSORPTION VITAMIN D STATUS AND BONE DISEASE
TΙ
     AFTER BOWEL RESECTION FOR CROHNS DISEASE.
     KELLY S; SELLIN J; MEREDITH S; SITRIN M; RABB J; ZULUTSKY M; ROSENBERG I H
ΑU
     UNIV. CHIC., CHICAGO, ILL. 60637, USA.
CS
     81ST ANNUAL MEETING OF THE AMERICAN GASTROENTEROLOGICAL ASSOCIATION, SALT
SO
     LAKE CITY, UTAH, USA, MAY 17-23, 1980. GASTROENTEROLOGY. (1980) 78 (5 PART
     2), 1193.
     CODEN: GASTAB. ISSN: 0016-5085.
DΤ
     Conference
     BR; OLD
FS
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English

LA

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General Biology - Symposia, Transactions and Proceedings of Conferences,
    Congresses, Review Annuals 00520
    Biochemical Studies - Vitamins 10063
    Biochemical Studies - Sterols and Steroids 10067
    Biochemical Studies - Minerals 10069
    Anatomy and Histology, General and Comparative - Surgery 11105
    Pathology, General and Miscellaneous - Inflammation and Inflammatory
    Disease 12508
    Pathology, General and Miscellaneous - Therapy
                                                     12512
    Metabolism - Minerals *13010
    Metabolism - Fat-Soluble Vitamins *13016
    Digestive System - General; Methods *14001
      Digestive System - Pathology *14006
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
    Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
    Hominidae 86215
BC
    Miscellaneous Descriptors
ΙT
       ABSTRACT HUMAN
     1406-16-2 (VITAMIN D)
RN
    7440-70-2 (CALCIUM)
L123 ANSWER 32 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
    1979:230971 BIOSIS
ΑN
    BA68:33475
DN
    CHANGES OF THE CALCIUM METABOLISM IN CROHNS DISEASE.
ΤI
ΑU
    KOCIAN J
    BUDEJOVICKA 800, 146 22 PRAHA 4, CZECH.
CS
    CESK GASTROENTEROL VYZ, (1979) 33 (1), 26-31.
SO
     CODEN: CKGAAM. ISSN: 0009-0565.
     BA; OLD
FS
    Czech
LA
     In a group of 21 patients with different stages of Crohn's
AB
     disease of the small intestine, a reduced dietary Ca intake was found in
     those in the acute stage of the disease, a slightly higher Ca intake in
     the chronically sick and a normal intake in patients after resection of
     the affected portion of the gut. The reduced Ca absorption, investigated
     by Ca absorption curves and by means of the isotope 47Ca on a whole-body
     counter is most marked in the acutely sick, less marked in the chronically
     sick and least in the groups with the resected gut. In impaired bone
    mineralization, the order of the 3 groups is the same. Mineralization is
     influenced by a reduced dietary Ca intake as well as reduced intestinal
     absorption of this element, obviously due to affection of the intestinal
     wall and impaired conversion of vitamin D into its
     active metabolites.
     Radiation - Radiation and Isotope Techniques 06504
CC
     Biochemical Studies - Vitamins 10063
     Biochemical Studies - Sterols and Steroids 10067
     Biochemical Studies - Minerals 10069
     Biophysics - Membrane Phenomena 10508
     Anatomy and Histology, General and Comparative - Surgery 11105
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease 12508
     Pathology, General and Miscellaneous - Therapy
                                                      12512
     Metabolism - Sterols and Steroids 13008
     Metabolism - Minerals *13010
     Metabolism - Fat-Soluble Vitamins 13016
     Nutrition - Malnutrition; Obesity *13203
     Nutrition - Minerals
                          *13206
     Nutrition - Fat-Soluble Vitamins 13208
     Digestive System - General; Methods 14001
     Digestive System - Physiology and Biochemistry *14004
       Digestive System - Pathology *14006
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Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology 18006
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
BC
    Hominidae 86215
IT
    Miscellaneous Descriptors
        HUMAN BONE MINERALIZATION INTESTINAL ABSORPTION VITAMIN
RN
     1406-16-2 (VITAMIN D)
     7440-70-2 (CALCIUM)
L123 ANSWER 33 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
    1979:80010 BIOSIS
AN
DN
    BR17:20010
    SERUM 25 HYDROXY VITAMIN D LEVELS IN CHILDREN AND
ΤI
    ADOLESCENTS WITH INFLAMMATORY BOWEL DISEASE.
     FLEISCHMAN A R; DAUM F; DINARI G; AIGES H; ROSEN J F
ΑU
    Pediatr. Res., (1978) 12 (4 PART 2), 364.
SO
    CODEN: PEREBL. ISSN: 0031-3998.
DT
    Conference
    BR: OLD
FS
    Unavailable
LA
    Methods, Materials and Apparatus, General - Photography 01012
CC
     Radiation - Radiation and Isotope Techniques 06504
     Biochemical Studies - Vitamins 10063
     Biochemical Studies - Proteins, Peptides and Amino Acids 10064
     Biochemical Studies - Sterols and Steroids 10067
     Biochemical Studies - Minerals 10069
     Enzymes - Physiological Studies *10808
    Anatomy and Histology, General and Comparative - Radiologic Anatomy 11106
     Pathology, General and Miscellaneous - Diagnostic 12504
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Metabolism - Sterols and Steroids *13008
     Metabolism - Proteins, Peptides and Amino Acids *13012
     Metabolism - Fat-Soluble Vitamins
     Nutrition - Malnutrition; Obesity *13203
       Digestive System - Pathology *14006
     Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies
     *15002
     Bones, Joints, Fasciae, Connective and Adipose Tissue - General; Methods
     18001
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Pharmacology - Digestive System *22014
     Toxicology - Pharmacological Toxicology Pediatrics *25000
                                               22504
     Hominidae 86215
BC
IT
    Miscellaneous Descriptors
        ABSTRACT HUMAN AZULFIDINE STEROIDS BONE GROWTH MAL ABSORPTION DRUG
        TREATMENT CALCIUM PHOSPHORUS ALKALINE PHOSPHATASE TRANS AMINASE
        GASTROINTESTINAL-DRUG
     599-79-1 (AZULFIDINE)
RN
     7440-70-2 (CALCIUM)
     7723-14-0 (PHOSPHORUS)
     9013-05-2 (PHOSPHATASE)
     9031-66-7 (TRANS AMINASE)
     19356-17-3Q, 21343-40-8Q, 64719-49-9Q (25 HYDROXY VITAMIN
     D)
L123 ANSWER 34 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
AN · 1978:72301 BIOSIS
     BR15:15801
DN
     BONE HISTOLOGY AND VITAMIN D STATUS IN CROHNS
ΤT
```

DISEASE ASSESSMENT OF VITAMIN D THERAPY.

```
DRISCOLL R; MEREDITH S; WAGONFELD J; ROSENBERG I
ΑU
     Gastroenterology, (1977) 72 (5 PT 2), A-28-1051.
SO
     CODEN: GASTAB. ISSN: 0016-5085.
DT
     Conference
     BR; OLD
FS
     Unavailable
LA
CC
     Comparative Biochemistry, General 10010
     Biochemical Studies - Vitamins 10063
     Biophysics - Molecular Properties and Macromolecules 10506
     Anatomy and Histology, General and Comparative - Microscopic and
     Ultramicroscopic Anatomy *11108
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease *12508
     Pathology, General and Miscellaneous - Therapy
                                                      12512
     Metabolism - Fat-Soluble Vitamins 13016
     Nutrition - Fat-Soluble Vitamins *13208
       Digestive System - Pathology *14006
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Pharmacology - Clinical Pharmacology
                                            22005
     Pharmacology - Digestive System *22014
     Immunology and Immunochemistry - Immunopathology, Tissue Immunology
     *34508
BC
     Hominidae 86215
TΤ
     Miscellaneous Descriptors
        ABSTRACT HUMAN METAB-DRUG
RN
     1406-16-2 (VITAMIN D)
L123 ANSWER 35 OF 35 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
AN
     1976:17841 BIOSIS
DN
     BR12:17841
     QUANTITATIVE ANALYSIS OF SKELETAL GROWTH DE MINERALIZATION AND
TТ
     VITAMIN D STATUS IN PATIENTS WITH INFLAMMATORY
     BOWEL DISEASE.
     WAGONFELD J B; GENANT H K; MALL J C; BOLT M; VANDER HORST J; ROSENBERG I H
ΑU
     Gastroenterology, (1975) 68 (4 PART 2), 1065.
SO
     CODEN: GASTAB. ISSN: 0016-5085.
DT
     Conference
FS
     BR; OLD
     Unavailable
LA
     Radiation - Radiation and Isotope Techniques 06504
CC
     Biochemical Methods - Minerals 10059
     Biochemical Studies - Vitamins 10063
     Biochemical Studies - Sterols and Steroids 10067
     Biochemical Studies - Minerals 10069
     Biophysics - General Biophysical Techniques 10504
     External Effects - Light and Darkness 10604
     Pathology, General and Miscellaneous - Inflammation and Inflammatory
     Disease 12508
     Pathology, General and Miscellaneous - Therapy
                                                      12512
     Metabolism - Minerals *13010
     Nutrition - Malnutrition; Obesity *13203
     Nutrition - Fat-Soluble Vitamins *13208
       Digestive System - Pathology *14006
     Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies
     15002
     Endocrine System - Adrenals 17004
     Bones, Joints, Fasciae, Connective and Adipose Tissue - Pathology *18006
     Pharmacology - Drug Metabolism; Metabolic Stimulators 22003
     Pharmacology - Clinical Pharmacology
                                           22005
     Pharmacology - Connective Tissue, Bone and Collagen - Acting Drugs 22012
     Pharmacology - Digestive System *22014
     Pharmacology - Endocrine System *22016
     Toxicology - Pharmacological Toxicology *22504
```

BC Hominidae 86215 Miscellaneous Descriptors IT ABSTRACT CORTICO STEROID THERAPY TOXICITY GRANULOMATOUS ILEO COLITIS ULCERATIVE COLITIS VITAMIN D DEFICIENCY SERUM CALCIUM LEVEL PHOTON ABSORPTIOMETRY INORGANIC PHOSPHATE CONCENTRATION RN 1406-16-2 (VITAMIN D) 7440-70-2 (CALCIUM) 14265-44-2 (PHOSPHATE) => fil wpix FILE 'WPIX' ENTERED AT 16:17:09 ON 14 SEP 2002 COPYRIGHT (C) 2002 THOMSON DERWENT <20020910/UP> FILE LAST UPDATED: 10 SEP 2002 200258 <200258/DW> MOST RECENT DERWENT UPDATE DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE >>> SLART (Simultaneous Left and Right Truncation) is now available in the /ABEX field. An additional search field /BIX is also provided which comprises both /BI and /ABEX <<< >>> The BATCH option for structure searches has been enabled in WPINDEX/WPIDS and WPIX <<< >>> PATENT IMAGES AVAILABLE FOR PRINT AND DISPLAY <<< >>> FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES, SEE http://www.derwent.com/dwpi/updates/dwpicov/index.html <<< >>> FOR A COPY OF THE DERWENT WORLD PATENTS INDEX STN USER GUIDE, PLEASE VISIT: http://www.stn-international.de/training\_center/patents/stn guide.pdf <<< >>> FOR INFORMATION ON ALL DERWENT WORLD PATENTS INDEX USER GUIDES, PLEASE VISIT: http://www.derwent.com/userguides/dwpi\_guide.html <<< => d all abeq tech abex tot L139 ANSWER 1 OF 4 WPIX (C) 2002 THOMSON DERWENT 2001-514277 [56] WPIX AN DNC C2001-153610 Use of vitamin D compounds for prevention and treatment of inflammatory bowel disease in humans and animals. DC B01 B05 HAYES, C E; NASHOLD, F E IN PA (NLIG-N) NORTHERN LIGHTS PHARM LLC CYC WO 2001046132 A1 20010628 (200156) \* EN 54p C07C401-00 PΤ RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW C07C401-00 AU 2001022878 A 20010703 (200164) US 6358939 B1 20020319 (200224) A61K031-593 WO 2001046132 A1 WO 2000-US34913 20001221; AU 2001022878 A AU 2001-22878

20001221; US 6358939 B1 US 1999-469985 19991221

FDT AU 2001022878 A Based on WO 200146132

```
PRAI US 1999-469985
                     19991221
    ICM A61K031-593; C07C401-00
    ICS A61K031-593
    WO 200146132 A UPAB: 20011001
AB
    NOVELTY - Vitamin D compounds or their compositions
    are administered to treat or prevent inflammatory bowel
    disease.
         ACTIVITY - Antiulcer; Antiinflammatory.
         C3H/HeJ strain mice were given DS (dextran sulfate (3.5 w/v%)) in
    acidified water for 5 days followed by acidified water without DS and
    continuously fed a synthetic diet. The mice showed no signs of
    colitis. The mice shunned the water containing DS and met their
    hydration means by consuming the synthetic diet. A control mice was given
    DS in acidified water, followed by acidified water without DS and
    continuously fed laboratory chow. The mice showed weight loss and had
    hemoglobin in the stool and thus the colitis was induced in the
    control mice.
         MECHANISM OF ACTION - Calcitriol inhibitor.
         C3H/HeJ strain mice were fed with a purified diet containing
    calcitriol (50 ng/day females; 200 ng/day males). On day 2, the mice were
    weighed and dextran sulfate (DS) (3.5 wt/vol%) was given in the drinking
    water on days 2 - 6. The mice were given acidified drinking water without
    DS for days 7 - 22. On days 7, 11, 15 and 19 mice were weighed and stool
    samples were collected. A blood sample was collected on 11 day. On day 22,
    mice were weighed, euthanized and stool, blood and colon samples were
    collected. A mock-treated control mice was also tested. The result showed
     that the calcitrol-treated mice exhibited significantly reduced weight
    loss, bloody diarrhea, shortening and thickening of the colon
    histopathologic score and inflammatory infiltration as compared
    to the mock-treated control.
         USE - For the prevention and treatment of inflammatory
    bowel disease e.g. Crohn's disease and
    ulcerative colitis in human, non-human primate, horse,
    dog or cat (preferably a mammal). The human is selected from a young adult
    living in united states, England, Northern Europe, Jewish descent,
    developing nation, a person with a family members who suffer from
    inflammatory bowel disease or a person
    determined to carry an IBD (inflammatory bowel
    disease) risk gene (all claimed).
         ADVANTAGE - The administration does not cause serious hypercalcemia.
    Administration delays onset symptoms of inflammatory
    bowel disease (all claimed). Vitamin D
     compounds can be administered in a cost-effective manner and timely
     fashion with a minimum of adverse side effects.
    Dwg.0/2
FS
    CPI
FA
    AB; DCN
    CPI: B10-E04A; B14-C03; B14-E08; B14-E10C; B14-L06
MC
                   UPTX: 20011001
TECH
    TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Compound: The
    vitamin D compound is selected from analogs of formula
     (I).
    X1 and X2 = H or acyl;
    Y1 and Y2 = H, O-aryl or O-alkyl having beta or alpha configuration;
    Z1 and Z2 = H;
    Z1+Z2 = CH2;
    R = Q or a group of formula (II);
    Q = alkyl, hydroxyalkyl or fluoroalkyl;
     a = S or R configuration;
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R1 = H, OH or O-acyl;

R2 and R3 = Q; R2+R3 = (CH2)m; m = 2 - 5;

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R4 = O-acyl or T;
     R5 = T;
     T = H, OH, F or Q;
     R4+R5 = double bonded O;
     R6+R7 = carbon-carbon double bond;
     R8 = H \text{ or } CH3;
    n = 1 - 5;
    X = CH, S, O or N.
     Provided that when Y1 is O-aryl or O-alkyl, Y2 is H and when Y1 is H, Y2
     is O-aryl or O-alkyl.
     Preferred Composition: The composition further comprises a transdermal
    patch.
ABEX
     SPECIFIC COMPOUNDS - Vitamin D, lalpha, 25-(OH) 2-16-ene-D3,
     lalpha, 25-(OH) 2-24-oxo16-ene-D3, lalpha, 24R(OH) 2-D3, lalpha, 25(OH) 2-22-oxa-
     D3, 20-epi-22-Oxa-24a,24b-dihomo-lalpha,25(OH)2-D3, 20-epi-22-oxa-
     24a,26a,27a-trihomo-lalpha,25(OH)2-D3, 20-epi-22-oxa-24homo-lalpha,25(OH)2-
     D3 and 1,25-(OH)2-16,23E-diene, 26-trifluoro-19-nor-D3 are specifically
     claimed as the vitamin D compounds.
     ADMINISTRATION - The route of administration can be intravenous, oral,
     parenteral, topical and rectal. The dosage is 0.1 - 20 microg per day per
     160 pound subject (all claimed).
     EXAMPLE - None given.
L139 ANSWER 2 OF 4 WPIX (C) 2002 THOMSON DERWENT
     2001-451613 [48]
                        WPIX
DNC C2001-136371
     Use of vitamin D compounds for treating or preventing
     inflammatory bowel disease, particularly
     ulcerative colitis or Crohn's disease.
     B01 B05
     CANTORNA, M T
     (PENN-N) PENN STATE RES FOUND
CYC
   94
    WO 2001042205 A2 20010614 (200148)* EN
                                              33p
                                                     C07C401-00
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
            NL OA PT SD SE SL SZ TR TZ UG ZW
         W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM
            DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC
            LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE
            SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW
     AU 2001045100 A 20010618 (200161)
                                                     C07C401-00
    WO 2001042205 A2 WO 2000-US42393 20001130; AU 2001045100 A AU 2001-45100
ADT
     20001130
    AU 2001045100 A Based on WO 200142205
PRAI US 2000-231906P 20000911; US 1999-168501P 19991202; US 2000-197827P
     20000414; US 2000-208632P 20000601
     ICM C07C401-00
     WO 200142205 A UPAB: 20010829
     NOVELTY - Use of vitamin D compounds for treating or
     preventing inflammatory bowel disease is
     new.
          ACTIVITY - Antiinflammatory.
          MECHANISM OF ACTION - T cell regulator.
          USE - For treating or preventing inflammatory bowel
     disease, particularly ulcerative colitis or
     Crohn's disease . The patient is on a low calcium diet (all
     claimed).
     Dwg.0/2
     CPI
```

AN

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PA

PΙ

IC

AB

FS

FA

AB; DCN

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CPI: B03-G; B05-A04; B14-C03; B14-E10C
MC
                   UPTX: 20010829
TECH
    TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Compounds: Preferred
    vitamin D compounds are of formula (I):
    Y1, Y2 = H or a hydroxy protecting group;
    Z1, Z2 = H; or together form =CH2;
    X1, X2 = H; or 1 is H and the other is O-aryl, O-alkyl, alkyl,
    hydroxyalkyl or fluoroalkyl; or together form =CR6R7;
    R6, R7 = H, alkyl, hydroxyalkyl or fluoroalkyl; or together form
     (CH2)x-;
    x = 2-5;
    R = a group of formula (i):
       = Y, -OY, CH2OY, -CCY or -CH=CHY;
    Y = H, Me, -COR5 or -(CH2)m C(R1)(R2)-(CH2)n-C(R3)(R4)(R5);
    m, n = 0-5;
    R1 = H, deuterium, OH, protected hydroxy, F, CF3 or 1-5C alkyl
    optionally substituted with hydroxy or protected hydroxy;
    R2, R3, R4 = deuterium, deuteroalkyl, H, F, CF3 or 1-5C alkyl optionally
    substituted with hydroxy or protected hydroxy; or
    R1+R2 may form oxo, = CR2R3 or -(CH2)p-; or R3+R4 may form oxo or
    -(CH2)q-;
    p, q = 2-5;
    R5 = H, 1-5C alkyl or optionally protected hydroxy;
    where any of the CH groups at positions 20, 22 or 23 in the side chain may
    be replaced by N, or any -CH(Me)-, -CH(R3)- or CH(R2)- at positions 20, 22
    and 23 respectively may be replaced by 0 or S.
ABEX
```

SPECIFIC COMPOUNDS - Preferred compounds include e.g. 1,25 dihydroxyvitamin D3.

ADMINISTRATION - Administration is oral, parenteral or transdermal. Daily dosage is 0.01--100~mug/day (all claimed).

EXAMPLE - 3 Week old vitamin D deficient wild-type (WT) and IL-10 knockout (KO) mice were either maintained vitamin D deficient or treated with cholecalciferol (5 microg/day). In a second series of experiments, 3 week old vitamin D deficient mice were maintained on the vitamin D deficient diet or supplemented with 1,25(OH)2D3 (0.005 microg/day), and sacrificed 4 weeks later. In a third series of experiments, 1,25(OH)2D3 treatment was started at the first signs of irritable bowel disease (IBD) (diarrhea, 7 weeks). 7 Week old vitamin D deficient mice were split into 2 groups; 1 group was maintained vitamin D deficient and the other was supplemented with 1,25(OH)2D3 (0.2 microg/day). Mice were treated for 2 weeks, then sacrificed.

There were no significant differences in the weight of any of the mice following 2 weeks treatment with 1,25 dihydroxycholecalciferol. However, the small intestines (SI) of the vitamin D deficient L-10 KO mice were enlarged and weighed significantly more than the SI from 1,25(OH)2D3 supplemented IL 10 KO, vitamin D deficient WT and 1,25(OH)2D3 supplemented WT mice. The SI from vitamin D deficient IL-10 KO mice were 9.9% of the total body weight, which is 2-fold higher than normal (about 5%). Treatment with 1,25(OH)2D3 for as little as 2 weeks reduced the inflammation in the SI of IL-10 KO mice.

```
AN 2001-353222 [37] WPIX
DNC C2001-109402
TI Multi-vitamin and mineral nutritional compositions for use in treating inflammatory bowel diseases including Crohn's disease, ulcerative colitis and celiac disease.
DC A96 B05 D13
IN SNOWDEN, R B
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L139 ANSWER 3 OF 4 WPIX (C) 2002 THOMSON DERWENT

```
(SNOW-N) SNOWDEN SUTTON ASSOC INC; (SNOW-I) SNOWDEN R B
PA
CYC 20
    US 6214373
                   B1 20010410 (200137)*
PΙ
                                                     A61K047-00
     WO 2001024642 Al 20010412 (200137) EN
                                                     A23K001-165
        RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE
    US 6214373 B1 US 1999-414666 19991007; WO 2001024642 A1 WO 2000-US27404
ADT
     20001005
PRAI US 1999-414666
                     19991007
    ICM A23K001-165; A61K047-00
          6214373 B UPAB: 20010704
AB
    NOVELTY - A nutritional composition for treating patients with
     inflammatory bowel diseases comprises selected
    proportions of multi-vitamins and minerals.
          DETAILED DESCRIPTION - A nutritional composition comprises: vitamin A
     (1,500-5,000 IU), vitamin D (200-600 IU), vitamin E
     (15-100 IU), vitamin K (15-60 mcg), vitamin C (30-150 mg), vitamin B1 (1-6
    mg), vitamin B2 (1-6 mg), vitamin B6 (1-6 mg), vitamin B12 (150-1,000
    meg), folic acid (0.2-0.5 mg), niacin (5-20 mg), biotin (0.1-0.2 mg),
    pantothenic acid (2-8 mg), iron (6-20 mg), calcium (50-200 mg), zinc (5-15
    mg), selenium (20-50 \text{ mcg}), copper (0.5-1.5 \text{ mg}), iodine
                                                              (60-80 mcg) and
    manganese (0.5-1.5 \text{ mg}). Wherein the minerals are included as salts other
     than carbonates.
          An INDEPENDENT CLAIM is also included for method for the treatment
     of inflammatory bowel disease or celiac
    disease.
          ACTIVITY - Antiinflammatory; antiulcer.
          No biological data given.
          MECHANISM OF ACTION - None given.
          USE - The nutritional composition is used for treating patients with
     inflammatory bowel diseases e.g. Crohn
     's disease, ulcerative colitis or celiac disease.
          ADVANTAGE - The composition is essentially free of magnesium which
     can act as a cathartic and free of carbonates which can generate gas in
     the gastrointestinal tract. The composition provides minerals and vitamins
     in a form and quantity which can help alleviate deficiencies which can be
     present in sufferers of inflammatory bowel
     diseases (IBD) e.g. Fe, Zn and vitamin C deficiencies are common
     in sufferers of IBD
     Dwg.0/0
FS
    CPI
    AB; DCN
FΑ
    CPI: A03-A00A; A12-V01; B03-A; B03-B; B03-C; B03-D; B03-E; B03-F;
MC.
          B03-G; B03-H; B03-J; B05-A01B; B05-A03; B05-C07; B14-C03;
          B14-E08; B14-E10; D03-H01T2
TECH
                    UPTX: 20010704
     TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Compositions: Mineral salts
     include phosphates, sulfates or fumarates. Iron is preferably present as
     ferrous fumarate and calcium as calcium diphosphate and the composition is
     free of magnesium. The composition may additionally comprise excipients
     selected from carboxymethylcellulose, microcrystalline cellulose, starch
     or modified starch. A particularly preferred composition comprises:
     vitamin A (2500 IU, retinyl acetate), vitamin D (400
     IU, cholecalciferol), vitamin E (75 IU, dl-alpha tocopherol acetate),
     vitamin k (40 mcg, phytonadione), vitamin C (100 mg, ascorbic acid),
     vitamin B1 (5 mg, thiamine mononitrate), vitamin B2 (5 mg, riboflavin),
     vitamin B6 (5 mg, pyridoxine hydrochloride), vitamin B12 (500 mcg,
     cyanocobalamin), folic acid (0.2 mg), niacin (10 mg, niacinamide),
     (0.15 mg), pantothenic acid (5 mg), iron (15 mg), calcium (100 mg),
     (11.25 mg), selenium (35 mcg), copper (1 mg), iodine (75 mcg) and
     manganese (1 mg).
ABEX
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ADMINISTRATION - Administration is oral as a unit dosage form e.g. a

tablet, caplet or capsule or in a liquid dosage form and administration is preferably twice daily (claimed).

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L139 ANSWER 4 OF 4 WPIX (C) 2002 THOMSON DERWENT
    1996-455225 [45]
                       WPIX
DNC
    C1996-142726
     Use of differentiating agents - for decreasing the inflammation associated
TI
     with chronic inflammatory intestinal conditions in patients.
DC
     B05 D16
    WU, G D
IN
     (UYPE-N) UNIV PENNSYLVANIA
PA
CYC 20
                   A1 19961003 (199645) * EN
                                              19p
                                                     C07C051-09
PΙ
    WO 9630326
        RW: AT BE CH DE DK ES FI FR GB GR IE IT LU MC NL PT SE
        W: CA JP
                   A 19991109 (199954)
                                                      A01N037-18
     US 5981597
    WO 9630326 A1 WO 1996-US4348 19960329; US 5981597 A CIP of US 1995-387116
ADT
     19950213, US 1995-413806 19950330
    US 5981597 A CIP of US 5569680
                    19950330; US 1995-387116
                                                 19950213
PRAI US 1995-413806
REP
    7.Jnl.Ref
     ICM A01N037-18; C07C051-09
IC
         A01N037-02; C08F022-14; C12N015-25
    ICS
          9630326 A UPAB: 19961111
AB
     A method is claimed for decreasing the inflammation associated with a
     chronic inflammatory intestinal condition in a patient comprising
     administering a differentiating agent, opt. in conjuncture with an
     inhibitor of inflammatory mediators produced by lymphocytes.
          USE - The method can be used to treat diseases such as
    ulcerative colitis, Crohn's disease, Type A or
     B chronic gastritis and graft vs. host diseases.
          ADVANTAGE - The differentiating agents alter the state of
     proliferation and ultimately the differentiation of colonic epithelial
     cells to reduce the inflammation. They also inhibit the expression of
     inflammatory mediators by epithelial cells.
     Dwg.0/2
FS
     CPI
FA
    AB; DCN
     CPI: B02-T; B03-A; B03-G; B04-H06F; B10-A10; B10-C04E; B10-D02;
MC.
          B10-E02; B10-G02; B14-C03; B14-E08; B14-E10B; B14-E10C; B14-E10D;
          D05-H
=> d his
     (FILE 'HOME' ENTERED AT 14:25:55 ON 14 SEP 2002)
                SET COST OFF
     FILE 'REGISTRY' ENTERED AT 14:26:21 ON 14 SEP 2002
                E VITAMIN D/CN
L1
              1 S E3
                STR
L2
             50 S L2 CSS
L3
     FILE 'HCAPLUS' ENTERED AT 14:28:01 ON 14 SEP 2002
                E HAYES C/AU
L4
             39 S E3, E5
                E HAYES COLEEN/AU
             52 S E4-E6
L5
                E NASHOLD F/AU
L6
             13 S E3-E6
            656 S (NORTH?(L)LIGHT?)/PA,CS
L7
            973 S (WISCON? (L) ALUM? (L) RES? (L) FOUND?) /PA, CS
L8
```

```
6480 S L1
L9
L10
          35010 S VITAMIN(S)D#
L11
           5664 S ?CALCIFERO?
             14 S L4, L5, L6 AND L9-L11
L12
            159 S L7, L8 AND L9-L11
L13
              5 S L12 AND L13
L14
              9 S L12 NOT L14
L15
           2619 S CALCITRIOL
L16
           2418 S 1 ALPHA 25 DIHYDROXYVITAMIN D3
L17
           5759 S 1 25 DIHYDROXYVITAMIN D3
L18
             78 S 1 ALPHA 25 DIHYDROXYVITAMIN D2
L19
             85 S 1 25 DIHYDROXYVITAMIN D2
L20
              9 S 19 NOR 1 ALPHA 25 DIHYDROXYVITAMIN D2
L21
              6 S 19 NOR 1 25 DIHYDROXYVITAMIN D2
L22
L23
             27 S PARICALCITOL
     FILE 'REGISTRY' ENTERED AT 14:35:26 ON 14 SEP 2002
              3 S 32222-06-3 OR 60133-18-8 OR 131918-61-1
L24
     FILE 'HCAPLUS' ENTERED AT 14:38:03 ON 14 SEP 2002
L25
           9086 S L24
             58 S ERCALCITRIOL OR ZEMPLAR OR RO176218 OR RO 17 6218 OR ROCALTRO
L26
           1399 S (1 25 OR 1 ALPHA 25)()(DIHYDROXYCALCIFEROL OR DIHYDROXYERGOCA
L27
           3791 S (1 25 OR 1 ALPHA 25)()OH 2D3
L28
             68 S (1 25 OR 1 ALPHA 25)()OH 2D2
L29
          30838 S ?VITAMIN? ()(D OR D2 OR D3)
L30
          36555 S ?VITAMIN? (S) (D OR D2 OR D3)
L31
L32
          42102 S L10, L11, L16-L23, L26-31
L33
          42200 S L32, L9, L25
     FILE 'REGISTRY' ENTERED AT 14:42:36 ON 14 SEP 2002
              9 S (32222-06-3 OR 60133-18-8 OR 131918-61-1)/CRN
L34
     FILE 'HCAPLUS' ENTERED AT 14:43:10 ON 14 SEP 2002
             14 S L5-L6 AND L33
L35
                SEL RN
     FILE 'REGISTRY' ENTERED AT 14:44:03 ON 14 SEP 2002
L36
             23 S E1-E23
              3 S L36 AND L1, L24
L37
L38
             20 S L36 NOT L37
             18 S L38 AND C5-C6/ES AND C6/ES
L39
                SEL RN 12 18 17
              3 S E24-E26
L40
L41
             15 S L39 NOT L40
                E 1.ALPHA., 25-DIHYDROXYVITAMIN D3/CN
L42
                E 19-NOR-1.ALPHA., 25-DIHYDROXYVITAMIN D2/CN
                E 1.ALPHA.-HYDROXYVITAMIN D3/CN
L43
              1 S E3
              1 S E2
L44
              3 S L1, L43, L44
L45
     FILE 'HCAPLUS' ENTERED AT 14:55:58 ON 14 SEP 2002
             15 S L21, L22
L46
     FILE 'REGISTRY' ENTERED AT 14:57:52 ON 14 SEP 2002
              1 S 131918-61-1
L47
              4 S L45, L47
L48
     FILE 'HCAPLUS' ENTERED AT 14:58:31 ON 14 SEP 2002
L49
           7460 S L48
             39 S PARICALCITOL OR ZEMPLAR OR L46
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L50

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78 S DOXERCALCIFEROL OR HECTOROL OR TSA840 OR TSA 840 OR 1()(HYDRO
L51
            130 S ALPHA CALCIDOL OR ALFACALCIDOL OR ALFAROL OR ALPHACALCIDOL OR
L52
             36 S 1() (HYDROXYCHOLECALCIFEROL OR HYDROXYVITAMIN D3 OR OH D3)
L53
            962 S 1()ALPHA() (HYDROXYCHOLECALCIFEROL OR HYDROXYVITAMIN D3 OR OH
L54
L55
          20797 S VITAMIN D OR CALCIFEROL
          21653 S L9, L49-L55
L56
L57
             13 S L4-L6 AND L56
                E INFLAMMATORY BOWEL/CT
                E E4+ALL
L58
           2993 S E2
                E INFLAMMATORY BOWEL/CT
                E E4+ALL
           3105 S INFLAMMATORY BOWEL() (DISEASE OR SYNDROME)
L59
           1077 S IBD
L60
                E ULCERATIVE COLITIS/CT
                E E3+ALL
           2115 S E2
L61
           3510 S ULCERATIVE ?COLITIS?
L62
                E CROHN/CT
                E E5+ALL
              0 S E2
L63
           1005 S CROHN?()(DISEASE OR ILEITIS OR INTESTIN? OR COLITIS)
L64
L65
             39 S L56 AND L58-L64
              1 S L57 AND L65
L66
             23 S L65 AND (PY<=1999 OR PRY<=1999 OR AY<=1999)
L67
             10 S (L49 OR L9) (L) (THU OR BAC OR USES) /RL AND L67
L68
                SEL DN AN 5 9
L69
              2 S E1-E6
                SEL DN AN L68 1-3
              3 S E7-E15
L70
              5 S L69,L70,L66 AND L4-L11,L16-L23,L25-L33,L35,L46,L49-L70
L71
                SEL RN L71 1
     FILE 'REGISTRY' ENTERED AT 15:37:03 ON 14 SEP 2002
             11 S E16-E26
L72
              1 S L72 AND L48
L73
             10 S L72 NOT L73
L74
              9 S L74 NOT CA
L75
     FILE 'HCAPLUS' ENTERED AT 15:37:45 ON 14 SEP 2002
                E DIGESTIVE TRACT/CT
                E E3+ALL
         141792 S E3, E101, E115
L76
            320 S E66, E68, E69, E72
L77
                E COLITIS/CT
                E E3+ALL
           3275 S E2
L78
                E INFLAMMATION/CT
           1308 S INFLAM?/CW (L) (INSTESTIN? OR BOWEL OR COLON? OR DIGEST? OR G
L79
T80
           2172 S L56 AND L76-L79
           2050 S L80 AND (PY<=1999 OR PRY<=1999 OR AY<=1999)
L81
             39 S L81 AND (CROHN? OR ?ULCER? OR BOWEL OR COLIT?)
L82
L83
             19 S L82 NOT L65
L84
              5 S L73, L75 AND L71
     FILE 'REGISTRY' ENTERED AT 15:44:58 ON 14 SEP 2002
     FILE 'HCAPLUS' ENTERED AT 15:45:27 ON 14 SEP 2002
     FILE 'MEDLINE' ENTERED AT 15:45:49 ON 14 SEP 2002
L85
           9879 S L48
L86
          22625 S L50-L55
L87
          22626 S L85, L86
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E INFLAMMATORY BOWEL/CT
                E E5+ALL
          29254 S E5+NT
L88
             77 S L87 AND L88
L89
             58 S L89 AND PY<=1999
L90
             11 S L90 NOT AB/FA
L91
L92
             47 S L90 NOT L91
             17 S L92 AND VITAMIN D/CT, CN
L93
             30 S L92 NOT L93
L94
                SEL DN AN 15
L95
              1 S L94 AND E1-E3
L96
              5 S (VITAMIN D) (L) TU/CT AND L93
              6 S L95, L96 AND L85-L96
L97
     FILE 'MEDLINE' ENTERED AT 15:54:30 ON 14 SEP 2002
     FILE 'EMBASE' ENTERED AT 15:54:40 ON 14 SEP 2002
L98
          22397 S L87
          18826 S L98 AND PY<=1999
L99
                E INFLAMMATORY BOWEL/CT
                E E5+ALL
                E E2+ALL
L100
          52398 S E12+NT
            105 S L99 AND L100
L101
             27 S L101 NOT AB/FA
L102
                SEL DN AN 6 22 26
L103
              3 S E1-E6
L104
             78 S L101 NOT L102
                E VITAMIN D/CT
          28988 S E3+NT
L105
             68 S L104 AND L105
L106
             16 S E3(L)DT/CT AND L106
L107
L108
              8 S L100 (L) DT/CT AND L107
L109
             11 S L103, L108
             70 S L104 NOT L109
L110
             11 S L109 AND L98-L110
L111
     FILE 'EMBASE' ENTERED AT 16:01:27 ON 14 SEP 2002
     FILE 'BIOSIS' ENTERED AT 16:01:37 ON 14 SEP 2002
          25213 S L87
L112
                E HAYES C/AU
L113
            202 S E3, E5
             24 S E48, E49
L114
                E NASHOLD F/AU
             14 S E3, E4
L115
             14 S L112 AND L113-L115
L116
L117
           1456 S 14006/CC AND L112
           1348 S *14006/CC AND L112
L118
             67 S L118 AND L64, L59, L62, L60
L119
L120
             34 S L119 AND PY<=1999
              1 S L116 AND L117
L121
             34 S L120 AND (?CROHN? OR ?INFLAM? OR ?COLIT? OR ?ULCER?)
L122
             35 S L121, L122
L123
     FILE 'BIOSIS' ENTERED AT 16:06:53 ON 14 SEP 2002
     FILE 'WPIX' ENTERED AT 16:07:08 ON 14 SEP 2002
L124
           1613 S L50-L55
                E VITAMIN D/DCN
                E E7+ALL
             55 S E2
L125
           1526 S (B03-G OR C03-G)/MC
L126
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L127	1854 S V340/M0,M1,M2,M3,M4,M5,M6
L128	2943 S L124-L127
L129	71 S L128 AND (?CROHN? OR ?INFLAM?(L)BOWEL OR ?COLIT? OR ?ULCER?)
L130	32 S L128 AND (CROHN? OR INFLAMMATORY BOWEL()(DISEASE OR SYNDROME)
	SEL DN AN 5 6 7
L131	3 S E1-E6
	SEL DN AN 28 L130
L132	1 S L130 AND E7-E8
L133	5 S L128 AND (HAYES C? OR NASHOLD F?)/AU
L134	1 S L128 AND (NORTH?(L)LIGHT?)/PA
L135	4 S L131,L132
L136	1 S L133,L134 AND L135
L137	4 S L135, L136
L138	4 S L133, L134 NOT L137
L139	4 S L137 AND L124-L138

FILE 'WPIX' ENTERED AT 16:17:09 ON 14 SEP 2002